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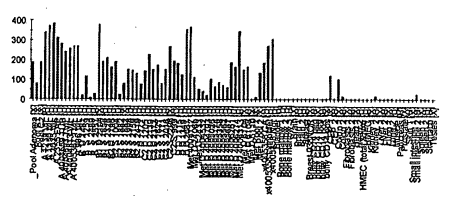


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(54) Title: NOVEL METHODS OF DIAGNOSING COLORECTAL CANCER, COMPOSITIONS, AND METHODS OF SCREENING FOR COLORECTAL CANCER MODULATORS



(57) Abstract

Described herein are methods that can be used for diagnosis and prognosis of colorectal cancer. Also described herein are methods that can be used to screen candidate bioactive agents for the ability to modulate colorectal cancer. Additionally, methods and molecular targets (genes and their products) for therapeutic intervention in colorectal and other cancers are described.

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NOVEL METHODS OF DIAGNOSING COLORECTAL CANCER, COMPOSITIONS, AND METHODS OF SCREENING FOR COLORECTAL CANCER MODULATORS

FIELD OF THE INVENTION

The invention relates to the identification of expression profiles and the nucleic acids involved in colorectal cancer, and to the use of such expression profiles and nucleic acids in diagnosis and prognosis of colorectal cancer. The invention further relates to methods for identifying and using candidate agents and/or targets which modulate colorectal cancer.

BACKGROUND OF THE INVENTION

Colorectal cancer is a significant cancer in Western populations. It develops as the result of a pathologic transformation of normal colon epithelium to an invasive cancer. There have been a number of recently characterized genetic alterations that have been implicated in colorectal cancer, including mutations in two classes of genes, tumor-suppressor genes and proto-oncogenes, with recent work suggesting that mutations in DNA repair genes may also be involved in tumorigenesis. For example, inactivating mutations of both alleles of the adenomatous polyposis coli (APC) gene, a tumor suppressor gene, appears to be one of the earliest events in colorectal cancer, and may even be the initiating event. Other genes implicated in colorectal cancer include the MCC gene, the p53 gene, the DCC (deleted in colorectal carcinoma) gene and other chromosome 18q genes, and genes in the TGF-β signaling pathway. For a review, see Molecular Biology of Colorectal Cancer, pp238-299, in Curr. Probl. Cancer, Sept/Oct 1997.

Imaging of colorectal cancer for diagnosis has been problematic and limited. In addition, dissemination of tumor cells (metastases) to locoregional lymph nodes is an important prognostic factor; five year survival rates drop from 80 percent in patients with no lymph node metastases to 45 to 50 percent in those patients who do have lymph node metastases. A recent report showed that micrometastases can be detected from lymph nodes using reverse transcriptase-PCR methods based on the presence of mRNA for carcinoembryonic antigen, which has previously been shown to be present in the vast

majority of colorectal cancers but not in normal tissues. Liefers et al., New England J. of Med. 339(4):223 (1998).

Thus, methods that can be used for diagnosis and prognosis of colorectal cancer would be desirable. Accordingly, provided herein are methods that can be used in diagnosis and prognosis of colorectal cancer. Further provided are methods that can be used to screen candidate bioactive agents for the ability to modulate colorectal cancer. Additionally, provided herein are molecular targets for therapeutic intervention in colorectal and other cancers.

SUMMARY OF THE INVENTION

The present invention provides methods for screening for compositions which modulate colorectal cancer. Also provided herein are methods of inhibiting proliferation of cell, preferably a colorectal cancer cell. Methods of treatment of cancer, as well as compositions, are also provided herein.

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In one aspect, a method of screening drug candidates comprises providing a cell that expresses an expression profile gene or fragments thereof. Preferred embodiments of the expression profile gene are genes which are differentially expressed in cancer cells, preferably colorectal cancer cells, compared to other cells. Preferred embodiments of expression profile genes used in the methods herein include but are not limited to the group consisting of CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7, CQA2, CGA8, CAA7 and CAA9; fragments of the proteins of this group are also preferred. It is understood that molecules for use in the present invention may be from any figure or any subset of listed molecules. Therefore, for example, any one or more of the genes listed above can be used in the methods herein. In another embodiment, a nucleic acid is selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14. Preferred nucleic acids are in Figure 12, more preferably Figure 13, and most preferably in Figure 14. The method further includes adding a drug candidate to the cell and determining the effect of the drug candidate on the expression of the expression profile gene.

In one embodiment, the method of screening drug candidates includes comparing the level of expression in the absence of the drug candidate to the level of expression in the presence of the drug candidate, wherein the concentration of the drug candidate can vary when present, and wherein the comparison can occur after addition or removal of the drug candidate. In a preferred embodiment, the cell expresses at least two expression profile genes. The profile genes may show an increase or decrease.

Also provided herein is a method of screening for a bioactive agent capable of binding to a colorectal cancer modulator protein (CCMP), the method comprising combining the CCMP and a candidate bioactive agent, and determining the binding of the candidate agent to the CCMP. Preferably the CCMP is a protein or fragment thereof selected from the group consisting of CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7, CQA2, CGA8, CAA7 and CAA9. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14. Preferred nucleic acids are in Figure 12, more preferably Figure 13, and most preferably in Figure 14.

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Further provided herein is a method for screening for a bioactive agent capable of modulating the activity of a CCMP. In one embodiment, the method comprises combining the CCMP and a candidate bioactive agent, and determining the effect of the candidate agent on the bioactivity of the CCMP. Preferably the CCMP is a protein or fragment thereof selected from the group consisting of CZA8. BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7, CQA2, CGA8, CAA7 and CAA9. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14. Preferred nucleic acids are in Figure 12, more preferably Figure 13, and most preferably in Figure 14.

Also provided is a method of evaluating the effect of a candidate colorectal cancer drug comprising administering the drug to a transgenic animal expressing or over-expressing the CCMP, or an animal lacking the CCMP, for example as a result of a gene knockout.

Additionally, provided herein is a method of evaluating the effect of a candidate colorectal cancer drug comprising administering the drug to a patient and removing a cell sample from the patient. The expression profile of the cell is then determined. This method may further comprise comparing the expression profile to an expression profile of a healthy individual.

Moreover, provided herein is a blochip comprising a nucleic acid segment which encodes a colorectal cancer protein, preferably selected from the group consisting of CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7, CQA2, CGA8, CAA7 and CAA9, or a fragment thereof, wherein the biochip comprises fewer than 1000 nucleic acid probes. Preferably at least two nucleic acid segments are included. In another embodiment, the nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14. Preferred nucleic acids are in Figure 12, more preferably Figure 13, and most preferably in Figure 14.

Furthermore, a method of diagnosing a disorder associated with colorectal cancer is provided. The method comprises determining the expression of a gene which encodes a colorectal cancer protein preferably selected from the group consisting of CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7, CQA2, CGA8, CAA7 and CAA9 or a fragment thereof in a first tissue type of a first individual, and comparing the distribution to the expression of the gene from a second normal tissue type from the first individual or a second unaffected individual. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14. Preferred nucleic acids are in Figure 12, more preferably Figure 13, and most preferably in Figure 14. A difference in the expression indicates that the first individual has a disorder associated with colorectal cancer.

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In another aspect, the present invention provides an antibody which specifically binds to a colorectal cancer protein, preferably selected from the group consisting of CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7, CQA2, CGA8, CAA7 and CAA9, or a fragment thereof. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14. Preferred nucleic acids are in Figure 12, more preferably Figure 13, and most preferably in Figure 14. In a preferred embodiment, the fragment of CAA9 is selected from CAA9p1, CAA9p2, CAA9p3, CAA9p4, CAA9p4MAPS, CAA9p5 and CAA9p5MAPS. Other preferred fragments for the breast cancer proteins are shown in the figures. Preferably the antibody is a monoclonal antibody. The antibody can be a fragment of an antibody such as a single stranded antibody as further described herein, or can be conjugated to another molecule. In one embodiment, the antibody is a humanized antibody.

In one embodiment a method for screening for a bioactive agent capable of interfering with the binding of a colorectal cancer protein (CCMP) or a fragment thereof and an antibody which binds to said CCMP or fragment thereof. In a preferred embodiment, the method comprises combining a CCMP or fragment thereof, a candidate bioactive agent and an antibody which binds to said CCMP or fragment thereof. The method further includes determining the binding of said CCMP or fragment thereof and said antibody. Wherein there is a change in binding, an agent is identified as an interfering agent. The interfering agent can be an agonist or an antagonist. Preferably, the antibody as well as the agent inhibits colorectal cancer.

In a further aspect, a method for inhibiting colorectal cancer is provided. In one embodiment, the method comprises administering to a cell a composition comprising an antibody to a colorectal modulating protein, preferably selected from the group consisting of CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7, CQA2, CGA8, CAA7 and CAA9, or a fragment

thereof. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14. Preferred nucleic acids are in Figure 12, more preferably Figure 13, and most preferably in Figure 14. The method can be performed in vitro or in vivo, preferably in vivo to an individual. In a preferred embodiment the method of inhibiting colorectal cancer is provided to an individual with cancer. As described herein, methods of inhibiting colorectal cancer can be performed by administering an inhibitor of colorectal cancer protein activity, including antisense molecules, and preferably small molecules.

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Also provided herein are methods eliciting an immune response in an individual. In one embodiment a method provided herein comprises administering to an individual a composition comprising a colorectal modulating protein, preferably selected from the group consisting of CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7, CQA2, CGA8, CAA7 and CAA9, or a fragment thereof. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14. Preferred nucleic acids are in Figure 12, more preferably Figure 13, and most preferably in Figure 14. In another aspect, said composition comprises a nucleic acid comprising a sequence encoding a colorectal cancer modulating protein, preferably selected from the group consisting of CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7, CQA2, CGA8, CAA7 and CAA9, or a fragment thereof. In another embodiment, the nucleic acid is selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14. Preferred nucleic acids are in Figure 12, more preferably Figure 13, and most preferably in Figure 14.

Further provided herein are compositions capable of eliciting an immune response in an individual. In one embodiment, a composition provided herein comprises a colorectal cancer modulating protein, preferably selected from the group consisting of CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7, CQA2, CGA8, CAA7 and CAA9, or a fragment thereof, and a pharmaceutically acceptable carrier. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14. Preferred nucleic acids are in Figure 12, more preferably Figure 13, and most preferably in Figure 14. In another embodiment, said composition comprises a nucleic acid comprising a sequence encoding a colorectal cancer modulating protein, preferably selected from the group consisting of CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7, CQA2, CGA8, CAA7 and CAA9, or a fragment thereof, and a pharmaceutically acceptable carrier. In another embodiment, the nucleic acid is selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14. Preferred nucleic acids are in Figure 12, more preferably Figure 13, and most preferably in Figure 14.

A method of neutralizing the effect of a colorectal cancer protein, preferably selected from the group consisting of CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7, CQA2, CGA8, CAA7 and CAA9, or a fragment thereof, comprising contacting an agent specific for said protein with said protein in an amount sufficient to effect neutralization. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14. Preferred nucleic acids are in Figure 12, more preferably Figure 13, and most preferably in Figure 14.

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In another aspect of the invention, a method of treating an individual for colorectal cancer is provided. In one embodiment, the method comprises administering to said individual an inhibitor of CJA8. In another embodiment, the method comprises administering to a patient having colorectal cancer an antibody to CJA8 conjugated to a therapeutic moiety. Such a therapeutic moiety can be a cytotoxic agent or a radioisotope.

Also provided herein is a method for determining the prognosis of an individual with colorectal cancer comprising determining the level of CJA8 in a sample, wherein a high level of CJA8 indicates a poor prognosis.

Novel sequences are also provided herein. Other aspects of the invention will become apparent to the skilled artisan by the following description of the invention.

DETAILED DESCRIPTION OF THE FIGURES

Figure 1 provides the Accession numbers for genes, including expression sequence tags, (incorporated in their entirety here and throughout the application where Accession numbers are provided), upregulated in tumor tissue compared to normal colon tissue.

Figure 2 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal colon tissue.

Figure 3 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal colon tissue.

Figure 4 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal colon tissue.

Figure 5 provides the Accession numbers for genes, including expression sequence tags, downregulated in tumor tissue compared to normal colon tissue.

Figure 6 provides the Accession numbers for genes, including expression sequence tags, downregulated in tumor tissue compared to normal colon tissue.

Figure 7 provides the Accession numbers for genes, including expression sequence tags, downregulated in tumor tissue compared to normal colon tissue.

Figure 8 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal colon tissue. Open reading frames in the sequences have been characterized as having a signal sequence (SS), a transmembrane domain (TM) or other.

Figure 9 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal colon tissue. Open reading frames in the sequences have been characterized as having a signal sequence (SS), a transmembrane domain (TM) or other.

Figure 10 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal colon tissue. Open reading frames have been characterized as having a signal sequence (SS), a transmembrane domain (TM) or other.

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Figure 11 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal colon tissue. Open reading frames have been characterized as having a signal sequence (SS), a transmembrane domain (TM) or other.

Figure 12 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal colon tissue. Open reading frames have been characterized as having a signal sequence (SS), a transmembrane domain (TM) or other.

Figure 13 provides the Accession numbers for genes or fragments thereof, including descriptions of the gene or encoded protein, upregulated in tumor tissue compared to normal colon tissue.

Figure 14 provides a list of proteins, including Accession numbers for nucleic acid sequences associated with the encoding genes thereof, upregulated in tumor tissue compared to normal colon tissue.

Figure 15 shows an embodiment of a nucleic acid which includes a sequence which encodes a colorectal protein provided herein, CAA2. The start and stop codon are shaded. The sequence within the two cross marks indicates a preferred novel fragment of CAA2 provided herein, referred to herein as the "CAA2 5' end". Preferred embodiments of CAA2 include at least a portion of the CAA2 5'. The sequence in bold and indicated with a bar at the bottom right beginning with "GGC" and ending with "AAA" can be found in Accession no. AA505133.

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Figure 16 shows an embodiment of a nucleic acid encoding CAA2, wherein the start and stop codons are shaded.

Figure 17 shows an embodiment of an amino acid sequence of CAA2. Preferred fragments include at least about 10 amino acids in the N-terminal end. The N-terminus as defined herein includes an embodiment beginning at the first amino acid until about any one of the three amino acids marked with a dot above them. In another embodiment, the N-terminus of CAA2 is defined as the amino acid sequence encoded by the CAA2 5' end.

Figure 18 shows the amino acid sequence of CAA2p1, a preferred CAA2 fragment provided herein.

Figure 19 shows the amino acid sequence of CAA2p2, a preferred CAA2 fragment provided herein.

Figure 20 shows an alignment of the human and mouse CAA2 polypeptides provided herein. The mouse polypeptide contains at least some of the sequence of each of the following Accession numbers: AA386837; Al508773; AA505293; and AA636546.

Figure 21 shows the relative amount of expression of CAA2 in various samples of colon cancer tissue 20 (dark bars) and many normal tissue types (light bars).

Figure 22 shows an embodiment of a colorectal cancer nucleic acid, CAA9 mRNA. The start and stop codons are underlined.

Figure 23 shows the open reading frame of the CAA9 gene wherein the start and stop codons are underlined.

Figure 24 shows an embodiment of the amino acid sequence of a colorectal cancer protein, CAA9, wherein putative transmembrane sequences are underlined. In one embodiment, CAA9 or fragments

of CAA9 are soluble, therefore, the transmembrane domains are deleted, inactivated, and/or the peptide is truncated (with or without re-ligation) to form soluble CAA9.

Figure 25 shows embodiments of colorectal cancer proteins (also termed colorectal cancer modulator proteins). Specifically, Figure 25 shows CAA9p1, CAA9p2, CAA9p3, CAA9p4, CAA9p4MAPS, CAA9p5 and CAA9p5MAPS and their respective solubilities.

Figure 26 shows the relative amount of CAA9 expression in several different samples of colon cancer tissue (dark bars) and normal tissues (light bars).

Figure 27 shows the nucleic acid sequence for the gene encoding CGA7. Start (ATG) and stop (TAG) codons are indicated by shaded boxes. In bold is the sequence of Accession No. AA331393.

Underlined is the consensus sequence derived from the compilation and alignment of published est sequences.

Figures 28A and 28B show the alignment summary and descriptions, respectively, of the various est's (by accession number) compiled to generate the consensus sequence of figure 1.

Figure 29 shows the amino acid sequence of CGA7.

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Figures 30A and 30B show the relative expression of CGA7 in normal tissue and colon cancer tissue, respectively.

Figure 31 shows the nucleic acid sequence for the mRNA encoding CGA8. Start (ATG) and stop (TAG) codons are indicated by shaded boxes. In bold is the sequence of Accession No. AA2786503. Underlined is the consensus sequence derived from the compilation and alignment of published est sequences.

Figures 32A and 32B show the alignment summary and descriptions, respectively, of the various est's (by accession number) compiled to generate the consensus sequence of figure 1.

Figure 33 shows the amino acid sequence of CGA8.

Figure 34 shows the relative expression of CGA8 in breast cancer tissue, colon cancer tissue, normal tissue and fetal tissue.

Figure 35 shows the sequence for the mRNA encoding CJA8. Start (ATG) and stop (TAA) codons are indicated by shaded boxes.

Figure 36 shows the amino acid sequence for CJA8. A putative transmembrane region is designated by shading. A mouse homolog of this human protein is found at Accession Number AAF21308.1.

Figure 37 shows the relative amount of expression of CJA8 in several different samples of colon tissues (dark bars) and normal tissues (light bars).

Figure 38 shows the relative amount of expression of BCN7 in several different samples of colon tissues (dark bars) and normal tissues (light bars), as determined using the sequence of Accession Number N22107 as a probe.

Figure 39 shows an embodiment of a nucleic acid which includes a sequence which encodes a colorectal cancer protein provided herein, BCN7.

Figure 40 shows the sequence for the mRNA encoding CZA8. Start (ATG) and stop (TGA) codons are indicated by underlining.

Figure 41 shows the sequence for the mRNA encoding BCX2. Start (ATG) and stop (TGA) codons are indicated by underlining.

Figure 42 shows the sequence for the mRNA encoding CBC2. Start (ATG) and stop (TAA) codons are indicated by underlining.

Figure 43 shows the sequence for the mRNA encoding CBC1. Start (ATG) and stop (TGA) codons are indicated by underlining.

Figure 44 shows the sequence for the mRNA encoding CBC3. Start (ATG) and stop (TGA) codons are indicated by underlining.

Figure 45 shows the sequence for the mRNA encoding BCN5. Start (ATG) and stop (TAA) codons are indicated by underlining.

Figure 46 shows an embodiment of a nucleic acid which includes a sequence which encodes a colorectal cancer protein provided herein, CJA9.

Figure 47 shows an embodiment of a nucleic acid which includes a sequence which encodes a colorectal cancer protein provided herein, CQA1.

Figure 48 shows an embodiment of a nucleic acid which includes a sequence which encodes a colorectal cancer protein provided herein, CQA2.

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DETAILED DESCRIPTION OF THE INVENTION

The present invention provides novel methods for diagnosis and prognosis evaluation for colorectal cancer (CRC), as well as methods for screening for compositions which modulate CRC. In one aspect, the expression levels of genes are determined in different patient samples for which either diagnosis or prognosis information is desired, to provide expression profiles. An expression profile of a particular sample is essentially a "fingerprint" of the state of the sample; while two states may have any particular gene similarly expressed, the evaluation of a number of genes simultaneously allows the generation of a gene expression profile that is unique to the state of the cell. That is, normal tissue may be distinguished from CRC tissue, and within CRC tissue, different prognosis states (good or poor long term survival prospects, for example) may be determined. By comparing expression profiles of colon tissue in known different states, information regarding which genes are important (including both up- and down-regulation of genes) in each of these states is obtained. The identification of sequences that are differentially expressed in CRC versus normal colon tissue, as well as differential expression resulting in different prognostic outcomes, allows the use of this information in a number of ways. For example, the evaluation of a particular treatment regime may be evaluated: does a chemotherapeutic drug act to improve the long-term prognosis in a particular patient. Similarly, diagnosis may be done or confirmed by comparing patient samples with the known expression profiles. Furthermore, these gene expression profiles (or individual genes) allow screening of drug candidates with an eye to mimicking or altering a particular expression profile; for example, screening can be done for drugs that suppress the CRC expression profile or convert a poor prognosis profile to a better prognosis profile. This may be done by making biochips comprising sets of the important CRC genes, which can then be used in these screens. These methods can also be done on the protein basis; that is, protein expression levels of the CRC proteins can be evaluated for diagnostic and prognostic purposes or to screen candidate agents. In addition, the CRC nucleic acid sequences can be administered for gene therapy purposes, including the administration of antisense nucleic acids, or the CRC proteins (including antibodies and other modulators thereof) administered as therapeutic drugs.

Thus the present invention provides nucleic acid and protein sequences that are differentially expressed in colorectal cancer, CRC, herein termed "CRC sequences". As outlined below, CRC sequences include those that are up-regulated (i.e. expressed at a higher level) in CRC, as well as those that are down-regulated (i.e. expressed at a lower level) in CRC. In a preferred embodiment, the CRC sequences are from humans; however, as will be appreciated by those in the art, CRC sequences from other organisms may be useful in animal models of disease and drug evaluation; thus, other CRC sequences are provided, from vertebrates, including mammals, including rodents (rats, mice, hamsters, guinea pigs, etc.), primates, farm animals (including sheep, goats, pigs, cows, horses, etc). CRC sequences from other organisms may be obtained using the techniques outlined below.

CRC sequences can include both nucleic acid and amino acid sequences. In a preferred embodiment, the CRC sequences are recombinant nucleic acids. By the term "recombinant nucleic acid" herein is meant nucleic acid, originally formed in vitro, in general, by the manipulation of nucleic acid by polymerases and endonucleases, in a form not normally found in nature. Thus an isolated nucleic acid, in a linear form, or an expression vector formed in vitro by ligating DNA molecules that are not normally joined, are both considered recombinant for the purposes of this invention. It is understood that once a recombinant nucleic acid is made and reintroduced into a host cell or organism, it will replicate non-recombinantly, i.e. using the in vivo cellular machinery of the host cell rather than in vitro manipulations; however, such nucleic acids, once produced recombinantly, although subsequently replicated non-recombinantly, are still considered recombinant for the purposes of the invention.

Similarly, a "recombinant protein" is a protein made using recombinant techniques, i.e. through the expression of a recombinant nucleic acid as depicted above. A recombinant protein is distinguished from naturally occurring protein by at least one or more characteristics. For example, the protein may be isolated or purified away from some or all of the proteins and compounds with which it is normally associated in its wild type host, and thus may be substantially pure. For example, an isolated protein is unaccompanied by at least some of the material with which it is normally associated in its natural state, preferably constituting at least about 0.5%, more preferably at least about 5% by weight of the total protein in a given sample. A substantially pure protein comprises at least about 75% by weight of the total protein, with at least about 80% being preferred, and at least about 90% being particularly preferred. The definition includes the production of a CRC protein from one organism in a different organism or host cell. Alternatively, the protein may be made at a significantly higher concentration than is normally seen, through the use of an inducible promoter or high expression promoter, such that the protein is made at increased concentration levels. Alternatively, the protein may be in a form

not normally found in nature, as in the addition of an epitope tag or amino acid substitutions, insertions and deletions, as discussed below.

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In a preferred embodiment, the CRC sequences are nucleic acids. As will be appreciated by those in the art and is more fully outlined below, CRC sequences are useful in a variety of applications, including diagnostic applications, which will detect naturally occurring nucleic acids, as well as screening applications; for example, biochips comprising nucleic acid probes to the CRC sequences can be generated. In the broadest sense, then, by "nucleic acid" or "oligonucleotide" or grammatical equivalents herein means at least two nucleotides covalently linked together. A nucleic acid of the present invention will generally contain phosphodiester bonds, although in some cases, as outlined below, nucleic acid analogs are included that may have alternate backbones, comprising, for example, phosphoramidate (Beaucage et al., Tetrahedron 49(10):1925 (1993) and references therein; Letsinger, J. Org. Chem. 35:3800 (1970); Sprinzl et al., Eur. J. Biochem. 81:579 (1977); Letsinger et al., Nucl. Acids Res. 14:3487 (1986); Sawai et al, Chem. Lett. 805 (1984), Letsinger et al., J. Am. Chem. Soc. 110:4470 (1988); and Pauwels et al., Chemica Scripta 26:141 91986)), phosphorothioate (Mag et al., Nucleic Acids Res. 19:1437 (1991); and U.S. Patent No. 5,644,048), phosphorodithioate (Briu et al., J. Am. Chem. Soc. 111:2321 (1989), O-methylphophoroamidite linkages (see Eckstein, Oligonucleotides and Analogues: A Practical Approach, Oxford University Press), and peptide nucleic acid backbones and linkages (see Egholm, J. Am. Chem. Soc. 114:1895 (1992); Meier et al., Chem. Int. Ed. Engl. 31:1008 (1992); Nielsen, Nature, 365:566 (1993); Carlsson et al., Nature 380:207 (1996), all of which are incorporated by reference). Other analog nucleic acids include those with positive backbones (Denpcy et al., Proc. Natl. Acad. Sci. USA 92:6097 (1995); non-lonic backbones (U.S. Patent Nos. 5,386,023, 5,637,684, 5,602,240, 5,216,141 and 4,469,863; Kiedrowshi et al., Angew. Chem. Intl. Ed. English 30:423 (1991); Letsinger et al., J. Am. Chem. Soc. 110:4470 (1988); Letsinger et al., Nucleoside & Nucleotide 13:1597 (1994); Chapters 2 and 3, ASC Symposium Series 580, "Carbohydrate Modifications in Antisense Research", Ed. Y.S. Sanghui and P. Dan Cook; Mesmaeker et al., Bioorganic & Medicinal Chem. Lett. 4:395 (1994); Jeffs et al., J. Biomolecular NMR 34:17 (1994); Tetrahedron Lett. 37:743 (1996)) and non-ribose backbones, including those described in U.S. Patent Nos. 5,235,033 and 5,034,506, and Chapters 6 and 7, ASC Symposium Series 580, "Carbohydrate Modifications in Antisense Research", Ed. Y.S. Sanghui and P. Dan Cook. Nucleic acids containing one or more carbocyclic sugars are also included within one definition of nucleic acids (see Jenkins et al., Chem. Soc. Rev. (1995) pp169-176). Several nucleic acid analogs are described in Rawls, C & E News June 2, 1997 page 35. All of these references are hereby expressly incorporated by reference. These modifications of the ribose-phosphate backbone may be done for a variety of reasons, for example to increase the stability and half-life of such molecules in physiological environments or as probes on a biochip.

As will be appreciated by those in the art, all of these nucleic acid analogs may find use in the present invention. In addition, mixtures of naturally occurring nucleic acids and analogs can be made; alternatively, mixtures of different nucleic acid analogs, and mixtures of naturally occurring nucleic acids and analogs may be made.

Particularly preferred are peptide nucleic acids (PNA) which includes peptide nucleic acid analogs. These backbones are substantially non-ionic under neutral conditions, in contrast to the highly charged phosphodiester backbone of naturally occurring nucleic acids. This results in two advantages. First, the PNA backbone exhibits improved hybridization kinetics. PNAs have larger changes in the melting temperature (Tm) for mismatched versus perfectly matched basepairs. DNA and RNA typically exhibit a 2-4°C drop in Tm for an internal mismatch. With the non-ionic PNA backbone, the drop is closer to 7-9°C. Similarly, due to their non-ionic nature, hybridization of the bases attached to these backbones is relatively insensitive to salt concentration. In addition, PNAs are not degraded by cellular enzymes, and thus can be more stable.

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The nucleic acids may be single stranded or double stranded, as specified, or contain portions of both double stranded or single stranded sequence. As will be appreciated by those in the art, the depiction of a single strand ("Watson") also defines the sequence of the other strand ("Crick"); thus the sequences described herein also includes the complement of the sequence. The nucleic acid may be DNA, both genomic and cDNA, RNA or a hybrid, where the nucleic acid contains any combination of deoxyribo- and ribo-nucleotides, and any combination of bases, including uracil, adenine, thymine, cytosine, guanine, inosine, xanthine hypoxanthine, isocytosine, isoguanine, etc. As used herein, the term "nucleoside" includes nucleotides and nucleoside and nucleotide analogs, and modified nucleosides such as amino modified nucleosides. In addition, "nucleoside" includes non-naturally occurring analog structures. Thus for example the individual units of a peptide nucleic acid, each containing a base, are referred to herein as a nucleoside.

A CRC sequence can be initially identified by substantial nucleic acid and/or amino acid sequence homology to the CRC sequences outlined herein. Such homology can be based upon the overall nucleic acid or amino acid sequence, and is generally determined as outlined below, using either homology programs or hybridization conditions.

The CRC sequences of the invention can be identified as follows. Samples of normal and tumor tissue are applied to biochips comprising nucleic acid probes. The samples are first microdissected, if applicable, and treated as is know in the art for the preparation of mRNA. Suitable biochips are

commercially available, for example from Affymetrix. Gene expression profiles as described herein are generated, and the data analyzed.

In a preferred embodiment, the genes showing changes in expression as between normal and disease states are compared to genes expressed in other normal tissues, including, but not limited to lung, heart, brain, liver, breast, kidney, muscle, prostate, small intestine, large intestine, spleen, bone, and placenta. In a preferred embodiment, those genes identified during the CRC screen that are expressed in any significant amount in other tissues are removed from the profile, although in some embodiments, this is not necessary. That is, when screening for drugs, it is preferable that the target be disease specific, to minimize possible side effects.

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In a preferred embodiment, CRC sequences are those that are up-regulated in CRC; that is, the expression of these genes is higher in colorectal carcinoma as compared to normal colon tissue. "Up-regulation" as used herein means at least about a two-fold change, preferably at least about a three fold change, with at least about five-fold or higher being preferred. All accession numbers herein are for the GenBank sequence database and the sequences of the accession numbers are hereby expressly incorporated by reference. GenBank is known in the art, see, e.g., Benson, DA, et al., Nucleic Acids Research 26:1-7 (1998) and http://www.ncbi.nlm.nih.gov/. In addition, these genes were found to be expressed in a limited amount or not at all in heart, brain, lung, liver, breast, kidney, prostate, small intestine and spleen.

In a preferred embodiment, CRC sequences are those that are down-regulated in CRC; that is, the expression of these genes is lower in colorectal carcinoma as compared to normal colon tissue. "Down-regulation" as used herein means at least about a two-fold change, preferably at least about a three fold change, with at least about five-fold or higher being preferred.

CRC proteins of the present invention may be classified as secreted proteins, transmembrane proteins or intracellular proteins. In a preferred embodiment the CRC protein is an intracellular protein. Intracellular proteins may be found in the cytoplasm and/or in the nucleus. Intracellular proteins are involved in all aspects of cellular function and replication (including, for example, signaling pathways); aberrant expression of such proteins results in unregulated or disregulated cellular processes. For example, many intracellular proteins have enzymatic activity such as protein kinase activity, protein phosphatase activity, protease activity, nucleotide cyclase activity, polymerase activity and the like. Intracellular proteins also serve as docking proteins that are involved in organizing complexes of proteins, or targeting proteins to various subcellular localizations, and are involved in maintaining the structural integrity of organelles.

An increasingly appreciated concept in characterizing intracellular proteins is the presence in the proteins of one or more motifs for which defined functions have been attributed. In addition to the highly conserved sequences found in the enzymatic domain of proteins, highly conserved sequences have been identified in proteins that are involved in protein-protein interaction. For example, Srchomology-2 (SH2) domains bind tyrosine-phosphorylated targets in a sequence dependent manner. PTB domains, which are distinct from SH2 domains, also bind tyrosine phosphorylated targets. SH3 domains bind to proline-rich targets. In addition, PH domains, tetratricopeptide repeats and WD domains to name only a few, have been shown to mediate protein-protein interactions. Some of these may also be involved in binding to phospholipids or other second messengers. As will be appreciated by one of ordinary skill in the art, these motifs can be identified on the basis of primary sequence; thus, an analysis of the sequence of proteins may provide insight into both the enzymatic potential of the molecule and/or molecules with which the protein may associate.

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In a preferred embodiment, the CRC sequences are transmembrane proteins. Transmembrane proteins are molecules that span the phospholipid bilayer of a cell. They may have an intracellular domain, an extracellular domain, or both. The intracellular domains of such proteins may have a number of functions including those already described for intracellular proteins. For example, the intracellular domain may have enzymatic activity and/or may serve as a binding site for additional proteins. Frequently the intracellular domain of transmembrane proteins serves both roles. For example certain receptor tyrosine kinases have both protein kinase activity and SH2 domains. In addition, autophosphorylation of tyrosines on the receptor molecule itself, creates binding sites for additional SH2 domain containing proteins.

Transmembrane proteins may contain from one to many transmembrane domains. For example, receptor tyrosine kinases, certain cytokine receptors, receptor guanylyl cyclases and receptor serine/threonine protein kinases contain a single transmembrane domain. However, various other proteins including channels and adenylyl cyclases contain numerous transmembrane domains. Many important cell surface receptors are classified as "seven transmembrane domain" proteins, as they contain 7 membrane spanning regions. Important transmembrane protein receptors include, but are not limited to insulin receptor, insulin-like growth factor receptor, human growth hormone receptor, glucose transporters, transferrin receptor, epidermal growth factor receptor, low density lipoprotein receptor, epidermal growth factor receptor, interleukin receptors, e.g. IL-1 receptor, IL-2 receptor, etc.

Characteristics of transmembrane domains include approximately 20 consecutive hydrophobic amino acids that may be followed by charged amino acids. Therefore, upon analysis of the amino acid

sequence of a particular protein, the localization and number of transmembrane domains within the protein may be predicted.

The extracellular domains of transmembrane proteins are diverse; however, conserved motifs are found repeatedly among various extracellular domains. Conserved structure and/or functions have been ascribed to different extracellular motifs. For example, cytokine receptors are characterized by a cluster of cysteines and a WSXWS (W= tryptophan, S= serine, X=any amino acid) motif.

Immunoglobulin-like domains are highly conserved. Mucin-like domains may be involved in cell adhesion and leucine-rich repeats participate in protein-protein interactions.

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Many extracellular domains are involved in binding to other molecules. In one aspect, extracellular domains are receptors. Factors that bind the receptor domain include circulating ligands, which may be peptides, proteins, or small molecules such as adenosine and the like. For example, growth factors such as EGF, FGF and PDGF are circulating growth factors that bind to their cognate receptors to initiate a variety of cellular responses. Other factors include cytokines, mitogenic factors, neurotrophic factors and the like. Extracellular domains also bind to cell-associated molecules. In this respect, they mediate cell-cell interactions. Cell-associated ligands can be tethered to the cell for example via a glycosylphosphatidylinositol (GPI) anchor, or may themselves be transmembrane proteins. Extracellular domains also associate with the extracellular matrix and contribute to the maintenance of the cell structure.

CRC proteins that are transmembrane are particularly preferred in the present invention as they are good targets for immunotherapeutics, as are described herein. In addition, as outlined below, transmembrane proteins can be also useful in imaging modalities.

It will also be appreciated by those in the art that a transmembrane protein can be made soluble by removing transmembrane sequences, for example through recombinant methods. Furthermore, transmembrane proteins that have been made soluble can be made to be secreted through recombinant means by adding an appropriate signal sequence.

In a preferred embodiment, the CRC proteins are secreted proteins; the secretion of which can be either constitutive or regulated. These proteins have a signal peptide or signal sequence that targets the molecule to the secretory pathway. Secreted proteins are involved in numerous physiological events; by virtue of their circulating nature, they serve to transmit signals to various other cell types. The secreted protein may function in an autocrine manner (acting on the cell that secreted the factor), a paracrine manner (acting on cells in close proximity to the cell that secreted the factor) or an

endocrine manner (acting on cells at a distance). Thus secreted molecules find use in modulating or altering numerous aspects of physiology. CRC proteins that are secreted proteins are particularly preferred in the present invention as they serve as good targets for diagnostic markers, for example for blood tests.

A CRC sequence is initially identified by substantial nucleic acid and/or amino acid sequence homology to the CRC sequences outlined herein. Such homology can be based upon the overall nucleic acid or amino acid sequence, and is generally determined as outlined below, using either homology programs or hybridization conditions.

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As used herein, a nucleic acid is a "CRC nucleic acid" if the overall homology of the nucleic acid sequence to the nucleic acid sequences encoding the amino acid sequences of the figures is preferably greater than about 75%, more preferably greater than about 80%, even more preferably greater than about 85% and most preferably greater than 90%. In some embodiments the homology will be as high as about 93 to 95 or 98%. Homology in this context means sequence similarity or identity, with identity being preferred. A preferred comparison for homology purposes is to compare the sequence containing sequencing errors to the correct sequence. This homology will be determined using standard techniques known in the art, including, but not limited to, the local homology algorithm of Smith & Waterman, Adv. Appl. Math. 2:482 (1981), by the homology alignment algorithm of Needleman & Wunsch, J. Mol. Biool. 48:443 (1970), by the search for similarity method of Pearson & Lipman, PNAS USA 85:2444 (1988), by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Drive, Madison, WI), the Best Fit sequence program described by Devereux et al., Nucl. Acid Res. 12:387-395 (1984), preferably using the default settings, or by inspection.

In a preferred embodiment, the sequences which are used to determine sequence identity or similarity are selected from the sequences set forth in the figures, preferably those represented in Figure 12, more preferably those represented in Figures 13A and 13B, still more preferably those of Figures 14-20, 22-25, 27-29, 31-33, 35-37 and 39-48, and fragments thereof. In one embodiment the sequences utilized herein are those set forth in the figures. In another embodiment, the sequences are naturally occurring allelic variants of the sequences set forth in the figures. In another embodiment, the sequences are sequence variants as further described herein.

One example of a useful algorithm is PILEUP. PILEUP creates a multiple sequence alignment from a group of related sequences using progressive, pairwise alignments. It can also plot a tree showing the

clustering relationships used to create the alignment. PILEUP uses a simplification of the progressive alignment method of Feng & Doolittle, J. Mol. Evol. 35:351-360 (1987); the method is similar to that described by Higgins & Sharp CABIOS 5:151-153 (1989). Useful PILEUP parameters including a default gap weight of 3.00, a default gap length weight of 0.10, and weighted end gaps.

5 Another example of a useful algorithm is the BLAST algorithm, described in Altschul et al., J. Mol. Biol. 215, 403-410, (1990) and Karlin et al., PNAS USA 90:5873-5787 (1993). A particularly useful BLAST program is the WU-BLAST-2 program which was obtained from Altschul et al., Methods in Enzymology, 266: 460-480 (1996); http://blast.wustl/edu/blast/ REACRCE.html]. WU-BLAST-2 uses several search parameters, most of which are set to the default values. The adjustable parameters 10 are set with the following values: overlap span =1, overlap fraction = 0.125, word threshold (T) = 11. The HSP S and HSP S2 parameters are dynamic values and are established by the program itself depending upon the composition of the particular sequence and composition of the particular database against which the sequence of interest is being searched; however, the values may be adjusted to increase sensitivity. A % amino acid sequence identity value is determined by the number 15 of matching identical residues divided by the total number of residues of the "longer" sequence in the aligned region. The "longer" sequence is the one having the most actual residues in the aligned region (gaps introduced by WU-Blast-2 to maximize the alignment score are ignored).

Thus, "percent (%) nucleic acid sequence identity" is defined as the percentage of nucleotide residues in a candidate sequence that are identical with the nucleotide residues of the sequences of the figures. A preferred method utilizes the BLASTN module of WU-BLAST-2 set to the default parameters, with overlap span and overlap fraction set to 1 and 0.125, respectively.

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The alignment may include the introduction of gaps in the sequences to be aligned. In addition, for sequences which contain either more or fewer nucleosides than those of the figures, it is understood that the percentage of homology will be determined based on the number of homologous nucleosides in relation to the total number of nucleosides. Thus, for example, homology of sequences shorter than those of the sequences identified herein and as discussed below, will be determined using the number of nucleosides in the shorter sequence.

In one embodiment, the nucleic acid homology is determined through hybridization studies. Thus, for example, nucleic acids which hybridize under high stringency to the nucleic acid sequences which encode the peptides identified in the figures, or their complements, are considered a CRC sequence. High stringency conditions are known in the art; see for example Maniatis et al., Molecular Cloning: A Laboratory Manual, 2d Edition, 1989, and Short Protocols in Molecular Biology, ed. Ausubel, et al.,

both of which are hereby incorporated by reference. Stringent conditions are sequence-dependent and will be different in different circumstances. Longer sequences hybridize specifically at higher temperatures. An extensive guide to the hybridization of nucleic acids is found in Tijssen, Techniques in Biochemistry and Molecular Biology--Hybridization with Nucleic Acid Probes, "Overview of principles of hybridization and the strategy of nucleic acid assays" (1993). Generally, stringent conditions are selected to be about 5-10°C lower than the thermal melting point (Tm) for the specific sequence at a defined ionic strength pH. The Tm is the temperature (under defined ionic strength, pH and nucleic acid concentration) at which 50% of the probes complementary to the target hybridize to the target sequence at equilibrium (as the target sequences are present in excess, at Tm, 50% of the probes are occupied at equilibrium). Stringent conditions will be those in which the salt concentration is less than about 1.0 M sodium ion, typically about 0.01 to 1.0 M sodium ion concentration (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes (e.g. 10 to 50 nucleotides) and at least about 60°C for long probes (e.g. greater than 50 nucleotides). Stringent conditions may also be achieved with the addition of destabilizing agents such as formamide.

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In another embodiment, less stringent hybridization conditions are used; for example, moderate or low stringency conditions may be used, as are known in the art; see Maniatis and Ausubel, supra, and Tijssen, supra.

In addition, the CRC nucleic acid sequences of the invention are fragments of larger genes, i.e. they are nucleic acid segments. "Genes" in this context includes coding regions, non-coding regions, and mixtures of coding and non-coding regions. Accordingly, as will be appreciated by those in the art, using the sequences provided herein, additional sequences of the CRC genes can be obtained, using techniques well known in the art for cloning either longer sequences or the full length sequences; see Maniatis et al., and Ausubel, et al., supra, hereby expressly incorporated by reference.

Once the CRC nucleic acid is identified, it can be cloned and, if necessary, its constituent parts recombined to form the entire CRC nucleic acid. Once isolated from its natural source, e.g., contained within a plasmid or other vector or excised therefrom as a linear nucleic acid segment, the recombinant CRC nucleic acid can be further-used as a probe to identify and isolate other CRC nucleic acids, for example additional coding regions. It can also be used as a "precursor" nucleic acid to make modified or variant CRC nucleic acids and proteins.

The CRC nucleic acids of the present invention are used in several ways. In a first embodiment, nucleic acid probes to the CRC nucleic acids are made and attached to biochips to be used in screening and diagnostic methods, as outlined below, or for administration, for example for gene

therapy and/or antisense applications. Alternatively, the CRC nucleic acids that include coding regions of CRC proteins can be put into expression vectors for the expression of CRC proteins, again either for screening purposes or for administration to a patient.

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In a preferred embodiment, nucleic acid probes to CRC nucleic acids (both the nucleic acid sequences encoding peptides outlined in the figures and/or the complements thereof) are made. The nucleic acid probes attached to the biochip are designed to be substantially complementary to the CRC nucleic acids, i.e. the target sequence (either the target sequence of the sample or to other probe sequences, for example in sandwich assays), such that hybridization of the target sequence and the probes of the present invention occurs. As outlined below, this complementarity need not be perfect; there may be any number of base pair mismatches which will interfere with hybridization between the target sequence and the single stranded nucleic acids of the present invention. However, if the number of mutations is so great that no hybridization can occur under even the least stringent of hybridization conditions, the sequence is not a complementary target sequence. Thus, by "substantially complementary" herein is meant that the probes are sufficiently complementary to the target sequences to hybridize under normal reaction conditions, particularly high stringency conditions, as outlined herein.

A nucleic acid probe is generally single stranded but can be partially single and partially double stranded. The strandedness of the probe is dictated by the structure, composition, and properties of the target sequence. In general, the nucleic acid probes range from about 8 to about 100 bases long, with from about 10 to about 80 bases being preferred, and from about 30 to about 50 bases being particularly preferred. That is, generally whole genes are not used. In some embodiments, much longer nucleic acids can be used, up to hundreds of bases.

In a preferred embodiment, more than one probe per sequence is used, with either overlapping probes or probes to different sections of the target being used. That is, two, three, four or more probes, with three being preferred, are used to build in a redundancy for a particular target. The probes can be overlapping (i.e. have some sequence in common), or separate.

As will be appreciated by those in the art, nucleic acids can be attached or immobilized to a solid support in a wide variety of ways. By "Immobilized" and grammatical equivalents herein is meant the association or binding between the nucleic acid probe and the solid support is sufficient to be stable under the conditions of binding, washing, analysis, and removal as outlined below. The binding can be covalent or non-covalent. By "non-covalent binding" and grammatical equivalents herein is meant one or more of either electrostatic, hydrophilic, and hydrophobic interactions. Included in non-covalent

binding is the covalent attachment of a molecule, such as, streptavidin to the support and the non-covalent binding of the biotinylated probe to the streptavidin. By "covalent binding" and grammatical equivalents herein is meant that the two moieties, the solid support and the probe, are attached by at least one bond, including sigma bonds, pi bonds and coordination bonds. Covalent bonds can be formed directly between the probe and the solid support or can be formed by a cross linker or by inclusion of a specific reactive group on either the solid support or the probe or both molecules. Immobilization may also involve a combination of covalent and non-covalent interactions.

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In general, the probes are attached to the biochip in a wide variety of ways, as will be appreciated by those in the art. As described herein, the nucleic acids can either be synthesized first, with subsequent attachment to the biochip, or can be directly synthesized on the biochip.

The biochip comprises a suitable solid substrate. By "substrate" or "solid support" or other grammatical equivalents herein is meant any material that can be modified to contain discrete individual sites appropriate for the attachment or association of the nucleic acid probes and is amenable to at least one detection method. As will be appreciated by those in the art, the number of possible substrates are very large, and include, but are not limited to, glass and modified or functionalized glass, plastics (including acrylics, polystyrene and copolymers of styrene and other materials, polypropylene, polyethylene, polybutylene, polyurethanes, TeflonJ, etc.), polysaccharides, nylon or nitrocellulose, resins, silica or silica-based materials including silicon and modified silicon, carbon, metals, inorganic glasses, plastics, etc. In general, the substrates allow optical detection and do not appreciably fluoresce. A preferred substrate is described in copending application entitled Reusable Low Fluorescent Plastic Biochip, U.S. Application Serial No. 09/270,214, filed March 15, 1999, herein incorporated by reference in its entirety.

Generally the substrate is planar, although as will be appreciated by those in the art, other configurations of substrates may be used as well. For example, the probes may be placed on the inside surface of a tube, for flow-through sample analysis to minimize sample volume. Similarly, the substrate may be flexible, such as a flexible foam, including closed cell foams made of particular plastics.

In a preferred embodiment, the surface of the biochip and the probe may be derivatized with chemical functional groups for subsequent attachment of the two. Thus, for example, the biochip is derivatized with a chemical functional group including, but not limited to, amino groups, carboxy groups, oxo groups and thiol groups, with amino groups being particularly preferred. Using these functional groups, the probes can be attached using functional groups on the probes. For example, nucleic

acids containing amino groups can be attached to surfaces comprising amino groups, for example using linkers as are known in the art; for example, homo-or hetero-bifunctional linkers as are well known (see 1994 Pierce Chemical Company catalog, technical section on cross-linkers, pages 155-200, incorporated herein by reference). In addition, in some cases, additional linkers, such as alkyl groups (including substituted and heteroalkyl groups) may be used.

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In this embodiment, the oligonucleotides are synthesized as is known in the art, and then attached to the surface of the solid support. As will be appreciated by those skilled in the art, either the 5' or 3' terminus may be attached to the solid support, or attachment may be via an internal nucleoside.

In an additional embodiment, the immobilization to the solid support may be very strong, yet non-covalent. For example, biotinylated oligonucleotides can be made, which bind to surfaces covalently coated with streptavidin, resulting in attachment.

Alternatively, the oligonucleotides may be synthesized on the surface, as is known in the art. For example, photoactivation techniques utilizing photopolymerization compounds and techniques are used. In a preferred embodiment, the nucleic acids can be synthesized in situ, using well known photolithographic techniques, such as those described in WO 95/25116; WO 95/35505; U.S. Patent Nos. 5,700,637 and 5,445,934; and references cited within, all of which are expressly incorporated by reference; these methods of attachment form the basis of the Affimetrix GeneChipTM technology.

In a preferred embodiment, CRC nucleic acids encoding CRC proteins are used to make a variety of expression vectors to express CRC proteins which can then be used in screening assays, as described below. The expression vectors may be either self-replicating extrachromosomal vectors or vectors which integrate into a host genome. Generally, these expression vectors include transcriptional and translational regulatory nucleic acid operably linked to the nucleic acid encoding the CRC protein. The term "control sequences" refers to DNA sequences necessary for the expression of an operably linked coding sequence in a particular host organism. The control sequences that are suitable for prokaryotes, for example, include a promoter, optionally an operator sequence, and a ribosome binding site. Eukaryotic cells are known to utilize promoters, polyadenylation signals, and enhancers.

Nucleic acid is "operably linked" when it is placed into a functional relationship with another nucleic acid sequence. For example, DNA for a presequence or secretory leader is operably linked to DNA for a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide; a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the

sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation. Generally, "operably linked" means that the DNA sequences being linked are contiguous, and, in the case of a secretory leader, contiguous and in reading phase. However, enhancers do not have to be contiguous. Linking is accomplished by ligation at convenient restriction sites. If such sites do not exist, the synthetic oligonucleotide adaptors or linkers are used in accordance with conventional practice. The transcriptional and translational regulatory nucleic acid will generally be appropriate to the host cell used to express the CRC protein; for example, transcriptional and translational regulatory nucleic acid sequences from *Bacillus* are preferably used to express the CRC protein in *Bacillus*. Numerous types of appropriate expression vectors, and suitable regulatory sequences are known in the art for a variety of host cells.

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In general, the transcriptional and translational regulatory sequences may include, but are not limited to, promoter sequences, ribosomal binding sites, transcriptional start and stop sequences, translational start and stop sequences, and enhancer or activator sequences. In a preferred embodiment, the regulatory sequences include a promoter and transcriptional start and stop sequences.

Promoter sequences encode either constitutive or inducible promoters. The promoters may be either naturally occurring promoters or hybrid promoters. Hybrid promoters, which combine elements of more than one promoter, are also known in the art, and are useful in the present invention.

In addition, the expression vector may comprise additional elements. For example, the expression vector may have two replication systems, thus allowing it to be maintained in two organisms, for example in mammalian or insect cells for expression and in a procaryotic host for cloning and amplification. Furthermore, for integrating expression vectors, the expression vector contains at least one sequence homologous to the host cell genome, and preferably two homologous sequences which flank the expression construct. The integrating vector may be directed to a specific locus in the host cell by selecting the appropriate homologous sequence for inclusion in the vector. Constructs for integrating vectors are well known in the art.

In addition, in a preferred embodiment, the expression vector contains a selectable marker gene to allow the selection of transformed host cells. Selection genes are well known in the art and will vary with the host cell used.

The CRC proteins of the present invention are produced by culturing a host cell transformed with an expression vector containing nucleic acid encoding a CRC protein, under the appropriate conditions to

induce or cause expression of the CRC protein. The conditions appropriate for CRC protein expression will vary with the choice of the expression vector and the host cell, and will be easily ascertained by one skilled in the art through routine experimentation. For example, the use of constitutive promoters in the expression vector will require optimizing the growth and proliferation of the host cell, while the use of an inducible promoter requires the appropriate growth conditions for induction. In addition, in some embodiments, the timing of the harvest is important. For example, the baculoviral systems used in insect cell expression are lytic viruses, and thus harvest time selection can be crucial for product yield.

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Appropriate host cells include yeast, bacteria, archaebacteria, fungi, and insect and animal cells, including mammalian cells. Of particular interest are *Drosophila melangaster* cells, *Saccharomyces* cerevisiae and other yeasts, *E. coli, Bacillus subtilis*, Sf9 cells, C129 cells, 293 cells, *Neurospora*, BHK, CHO, COS, HeLa cells, THP1 cell line (a macrophage cell line) and human cells and cell lines.

In a preferred embodiment, the CRC proteins are expressed in mammalian cells. Mammalian expression systems are also known in the art, and include retroviral systems. A preferred expression vector system is a retroviral vector system such as is generally described in PCT/US97/01019 and PCT/US97/01048, both of which are hereby expressly incorporated by reference. Of particular use as mammalian promoters are the promoters from mammalian viral genes, since the viral genes are often highly expressed and have a broad host range. Examples include the SV40 early promoter, mouse mammary tumor virus LTR promoter, adenovirus major late promoter, herpes simplex virus promoter, and the CMV promoter. Typically, transcription termination and polyadenylation sequences recognized by mammalian cells are regulatory regions located 3' to the translation stop codon and thus, together with the promoter elements, flank the coding sequence. Examples of transcription terminator and polyadenlytion signals include those derived form SV40.

The methods of introducing exogenous nucleic acid into mammalian hosts, as well as other hosts, is well known in the art, and will vary with the host cell used. Techniques include dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, viral infection, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei.

In a preferred embodiment, CRC proteins are expressed in bacterial systems. Bacterial expression systems are well known in the art. Promoters from bacteriophage may also be used and are known in the art. In addition, synthetic promoters and hybrid promoters are also useful; for example, the tac promoter is a hybrid of the trp and lac promoter sequences. Furthermore, a bacterial promoter can

include naturally occurring promoters of non-bacterial origin that have the ability to bind bacterial RNA polymerase and initiate transcription. In addition to a functioning promoter sequence, an efficient ribosome binding site is desirable. The expression vector may also include a signal peptide sequence that provides for secretion of the CRC protein in bacteria. The protein is either secreted into the growth media (gram-positive bacteria) or into the periplasmic space, located between the inner and outer membrane of the cell (gram-negative bacteria). The bacterial expression vector may also include a selectable marker gene to allow for the selection of bacterial strains that have been transformed. Suitable selection genes include genes which render the bacteria resistant to drugs such as ampicillin, chloramphenicol, erythromycin, kanamycin, neomycin and tetracycline. Selectable markers also include biosynthetic genes, such as those in the histidine, tryptophan and leucine biosynthetic pathways. These components are assembled into expression vectors. Expression vectors for bacteria are well known in the art, and include vectors for Bacillus subtilis, E. coli, Streptococcus cremoris, and Streptococcus lividans, among others. The bacterial expression vectors are transformed into bacterial host cells using techniques well known in the art, such as calcium chloride treatment, electroporation, and others.

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In one embodiment, CRC proteins are produced in insect cells. Expression vectors for the transformation of insect cells, and in particular, baculovirus-based expression vectors, are well known in the art.

In a preferred embodiment, CRC protein is produced in yeast cells. Yeast expression systems are well known in the art, and include expression vectors for Saccharomyces cerevisiae, Candida albicans and C. maltosa, Hansenula polymorpha, Kluyveromyces fragilis and K. lactis, Pichia guillerimondii and P. pastoris, Schizosaccharomyces pombe, and Yarrowia lipolytica.

The CRC protein may also be made as a fusion protein, using techniques well known in the art. Thus, for example, for the creation of monoclonal antibodies, if the desired epitope is small, the CRC protein may be fused to a carrier protein to form an immunogen. Alternatively, the CRC protein may be made as a fusion protein to increase expression, or for other reasons. For example, when the CRC protein is a CRC peptide, the nucleic acid encoding the peptide may be linked to other nucleic acid for expression purposes.

In one embodiment, the CRC nucleic acids, proteins and antibodies of the invention are labeled. By "labeled" herein is meant that a compound has at least one element, isotope or chemical compound attached to enable the detection of the compound. In general, labels fall into three classes: a) isotopic labels, which may be radioactive or heavy isotopes; b) immune labels, which may be antibodies or

antigens; and c) colored or fluorescent dyes. The labels may be incorporated into the CRC nucleic acids, proteins and antibodies at any position. For example, the label should be capable of producing, either directly or indirectly, a detectable signal. The detectable moiety may be a radioisotope, such as ³H, ¹⁴C, ³²P, ³⁵S, or ¹²⁵I, a fluorescent or chemiluminescent compound, such as fluorescein isothiocyanate, rhodamine, or luciferin, or an enzyme, such as alkaline phosphatase, betagalactosidase or horseradish peroxidase. Any method known in the art for conjugating the antibody to the label may be employed, including those methods described by Hunter et al., Nature, 144:945 (1962); David et al., Biochemistry, 13:1014 (1974); Pain et al., J. Immunol. Meth., 40:219 (1981); and Nygren, J. Histochem. and Cytochem., 30:407 (1982).

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Accordingly, the present invention also provides CRC protein sequences. A CRC protein of the present invention may be identified in several ways. "Protein" in this sense includes proteins, polypeptides, and peptides. As will be appreciated by those in the art, the nucleic acid sequences of the invention can be used to generate protein sequences. There are a variety of ways to do this, including cloning the entire gene and verifying its frame and amino acid sequence, or by comparing it to known sequences to search for homology to provide a frame, assuming the CRC protein has homology to some protein in the database being used. Generally, the nucleic acid sequences are input into a program that will search all three frames for homology. This is done in a preferred embodiment using the following NCBI Advanced BLAST parameters. The program is blastx or blastn. The database is nr. The input data is as "Sequence in FASTA format". The organism list is "none". The "expect" is 10; the filter is default. The "descriptions" is 500, the "alignments" is 500, and the "alignment view" is pairwise. The "Query Genetic Codes" is standard (1). The matrix is BLOSUM62; gap existence cost is 11, per residue gap cost is 1; and the lambda ratio is .85 default. This results in the generation of a putative protein sequence.

Also included within one embodiment of CRC proteins are amino acid variants of the naturally occurring sequences, as determined herein. Preferably, the variants are preferably greater than about 75% homologous to the wild-type sequence, more preferably greater than about 80%, even more preferably greater than about 85% and most preferably greater than 90%. In some embodiments the homology will be as high as about 93 to 95 or 98%. As for nucleic acids, homology in this context means sequence similarity or identity, with identity being preferred. This homology will be determined using standard techniques known in the art as are outlined above for the nucleic acid homologies.

CRC proteins of the present invention may be shorter or longer than the wild type amino acid sequences. Thus, in a preferred embodiment, included within the definition of CRC proteins are portions or fragments of the wild type sequences, herein. In addition, as outlined above, the CRC

nucleic acids of the invention may be used to obtain additional coding regions, and thus additional protein sequence, using techniques known in the art.

In a preferred embodiment, the CRC proteins are derivative or variant CRC proteins as compared to the wild-type sequence. That is, as outlined more fully below, the derivative CRC peptide will contain at least one amino acid substitution, deletion or insertion, with amino acid substitutions being particularly preferred. The amino acid substitution, insertion or deletion may occur at any residue within the CRC peptide.

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Also included in an embodiment of CRC proteins of the present invention are amino acid sequence variants. These variants fall into one or more of three classes: substitutional, insertional or deletional variants. These variants ordinarily are prepared by site specific mutagenesis of nucleotides in the DNA encoding the CRC protein, using cassette or PCR mutagenesis or other techniques well known in the art, to produce DNA encoding the variant, and thereafter expressing the DNA in recombinant cell culture as outlined above. However, variant CRC protein fragments having up to about 100-150 residues may be prepared by in vitro synthesis using established techniques. Amino acid sequence variants are characterized by the predetermined nature of the variation, a feature that sets them apart from naturally occurring allelic or interspecies variation of the CRC protein amino acid sequence. The variants typically exhibit the same qualitative biological activity as the naturally occurring analogue, although variants can also be selected which have modified characteristics as will be more fully outlined below.

While the site or region for introducing an amino acid sequence variation is predetermined, the mutation per se need not be predetermined. For example, in order to optimize the performance of a mutation at a given site, random mutagenesis may be conducted at the target codon or region and the expressed CRC variants screened for the optimal combination of desired activity. Techniques for making substitution mutations at predetermined sites in DNA having a known sequence are well known, for example, M13 primer mutagenesis and PCR mutagenesis. Screening of the mutants is done using assays of CRC protein activities.

Amino acid substitutions are typically of single residues; insertions usually will be on the order of from about 1 to 20 amino acids, although considerably larger insertions may be tolerated. Deletions range from about 1 to about 20 residues, although in some cases deletions may be much larger.

Substitutions, deletions, insertions or any combination thereof may be used to arrive at a final derivative. Generally these changes are done on a few amino acids to minimize the alteration of the

molecule. However, larger changes may be tolerated in certain circumstances. When small alterations in the characteristics of the CRC protein are desired, substitutions are generally made in accordance with the following chart:

		Chart I			
5	Original Residue	Exemplary Substitutions			
	Ala	Ser			
	Arg	Lys			
	Asn	Gln, His			
	Asp ·	Glu			
10	Cys	Ser			
	Gln ⁵	Asn			
	Glu	Asp			
	Gly	Pro			
	His	Asn, Gln			
15	, ile	Leu, Val			
	Leu	lle, Val			
	Lys	Arg, Gln, Glu			
	Met	Leu, lie			
	Phe	Met, Leu, Tyr			
20	Ser	Thr			
	Thr	Ser			
	Trp	Tyr			
	Tyr	Trp, Phe			
	Val	lle, Leu			

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Substantial changes in function or immunological identity are made by selecting substitutions that are less conservative than those shown in Chart I. For example, substitutions may be made which more significantly affect: the structure of the polypeptide backbone in the area of the alteration, for example the alpha-helical or beta-sheet structure; the charge or hydrophobicity of the molecule at the target site; or the bulk of the side chain. The substitutions which in general are expected to produce the greatest changes in the polypeptide's properties are those in which (a) a hydrophilic residue, e.g. seryl or threonyl is substituted for (or by) a hydrophobic residue, e.g. leucyl, isoleucyl, phenylalanyl, valyl or alanyl; (b) a cysteine or proline is substituted for (or by) any other residue; (c) a residue having an electropositive side chain, e.g. lysyl, arginyl, or histidyl, is substituted for (or by) an electronegative residue, e.g. glutamyl or aspartyl; or (d) a residue having a bulky side chain, e.g. phenylalanine, is substituted for (or by) one not having a side chain, e.g. glycine.

The variants typically exhibit the same qualitative biological activity and will elicit the same immune response as the naturally-occurring analogue, although variants also are selected to modify the characteristics of the CRC proteins as needed. Alternatively, the variant may be designed such that the biological activity of the CRC protein is altered. For example, glycosylation sites may be altered or removed.

Covalent modifications of CRC polypeptides are included within the scope of this invention. One type of covalent modification includes reacting targeted amino acid residues of a CRC polypeptide with an organic derivatizing agent that is capable of reacting with selected side chains or the N-or C-terminal residues of a CRC polypeptide. Derivatization with bifunctional agents is useful, for instance, for crosslinking CRC to a water-insoluble support matrix or surface for use in the method for purifying anti-CRC antibodies or screening assays, as is more fully described below. Commonly used crosslinking agents include, e.g., 1,1-bis(diazoacetyl)-2-phenylethane, glutaraldehyde, N-hydroxy-succinimide esters, for example, esters with 4-azidosalicylic acid, homobifunctional imidoesters, including disuccinimidyl esters such as 3,3'-dithiobis(succinimidylpropionate), bifunctional maleimides such as bis-N-maleimido-1,8-octane and agents such as methyl-3-[(p-azidophenyl)dithio]propioimidate.

Other modifications include deamidation of glutaminyl and asparaginyl residues to the corresponding glutamyl and aspartyl residues, respectively, hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of seryl, threonyl or tyrosyl residues, methylation of the α -amino groups of lysine, arginine, and histidine side chains [T.E. Creighton, Proteins: Structure and Molecular Properties, W.H. Freeman & Co., San Francisco, pp. 79-86 (1983)], acetylation of the N-terminal amine, and amidation of any C-terminal carboxyl group.

Another type of covalent modification of the CRC polypeptide included within the scope of this invention comprises altering the native glycosylation pattern of the polypeptide. "Altering the native glycosylation pattern" is intended for purposes herein to mean deleting one or more carbohydrate moleties found in native sequence CRC polypeptide, and/or adding one or more glycosylation sites that are not present in the native sequence CRC polypeptide.

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Addition of glycosylation sites to CRC polypeptides may be accomplished by altering the amino acid sequence thereof. The alteration may be made, for example, by the addition of, or substitution by, one or more serine or threonine residues to the native sequence CRC polypeptide (for O-linked glycosylation sites). The CRC amino acid sequence may optionally be altered through changes at the DNA level, particularly by mutating the DNA encoding the CRC polypeptide at preselected bases such that codons are generated that will translate into the desired amino acids.

Another means of increasing the number of carbohydrate moieties on the CRC polypeptide is by chemical or enzymatic coupling of glycosides to the polypeptide. Such methods are described in the art, e.g., in WO 87/05330 published 11 September 1987, and in Aplin and Wriston, CRC Crit. Rev. Biochem., pp. 259-306 (1981).

Removal of carbohydrate moieties present on the CRC polypeptide may be accomplished chemically or enzymatically or by mutational substitution of codons encoding for amino acid residues that serve as targets for glycosylation. Chemical deglycosylation techniques are known in the art and described, for instance, by Hakimuddin, et al., Arch. Biochem. Biophys., 259:52 (1987) and by Edge et al., Anal. Biochem., 118:131 (1981). Enzymatic cleavage of carbohydrate moieties on polypeptides can be achieved by the use of a variety of endo-and exo-glycosidases as described by Thotakura et al., Meth. Enzymol., 138:350 (1987).

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Another type of covalent modification of CRC comprises linking the CRC polypeptide to one of a variety of nonproteinaceous polymers, e.g., polyethylene glycol, polypropylene glycol, or polyoxyalkylenes, in the manner set forth in U.S. Patent Nos. 4,640,835; 4,496,689; 4,301,144; 4,670,417; 4,791,192 or 4,179,337.

CRC polypeptides of the present invention may also be modified in a way to form chimeric molecules comprising a CRC polypeptide fused to another, heterologous polypeptide or amino acid sequence. In one embodiment, such a chimeric molecule comprises a fusion of a CRC polypeptide with a tag polypeptide which provides an epitope to which an anti-tag antibody can selectively bind. The epitope tag is generally placed at the amino-or carboxyl-terminus of the CRC polypeptide. The presence of such epitope-tagged forms of a CRC polypeptide can be detected using an antibody against the tag polypeptide. Also, provision of the epitope tag enables the CRC polypeptide to be readily purified by affinity purification using an anti-tag antibody or another type of affinity matrix that binds to the epitope tag. In an alternative embodiment, the chimeric molecule may comprise a fusion of a CRC polypeptide with an immunoglobulin or a particular region of an immunoglobulin. For a bivalent form of the chimeric molecule, such a fusion could be to the Fc region of an IgG molecule.

Various tag polypeptides and their respective antibodies are well known in the art. Examples include poly-histidine (poly-his) or poly-histidine-glycine (poly-his-gly) tags; the flu HA tag polypeptide and its antibody 12CA5 [Field et al., Mol. Cell. Biol., 8:2159-2165 (1988)]; the c-myc tag and the 8F9, 3C7, 6E10, G4, B7 and 9E10 antibodies thereto [Evan et al., Molecular and Cellular Biology, 5:3610-3616 (1985)]; and the Herpes Simplex virus glycoprotein D (gD) tag and its antibody [Paborsky et al., Protein Engineering, 3(6):547-553 (1990)]. Other tag polypeptides include the Flag-peptide [Hopp et al., BioTechnology, 6:1204-1210 (1988)]; the KT3 epitope peptide [Martin et al., Science, 255:192-194 (1992)]; tubulin epitope peptide [Skinner et al., J. Biol. Chem., 266:15163-15166 (1991)]; and the T7 gene 10 protein peptide tag [Lutz-Freyermuth et al., Proc. Natl. Acad. Sci. USA, 87:6393-6397 (1990)].

Also included with the definition of CRC protein in one embodiment are other CRC proteins of the CRC family, and CRC proteins from other organisms, which are cloned and expressed as outlined below. Thus, probe or degenerate polymerase chain reaction (PCR) primer sequences may be used to find other related CRC proteins from humans or other organisms. As will be appreciated by those in the art, particularly useful probe and/or PCR primer sequences include the unique areas of the CRC nucleic acid sequence. As is generally known in the art, preferred PCR primers are from about 15 to about 35 nucleotides in length, with from about 20 to about 30 being preferred, and may contain inosine as needed. The conditions for the PCR reaction are well known in the art.

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In addition, as is outlined herein, CRC proteins can be made that are longer than those depicted in the figures, for example, by the elucidation of additional sequences, the addition of epitope or purification tags, the addition of other fusion sequences, etc.

CRC proteins may also be identified as being encoded by CRC nucleic acids. Thus, CRC proteins are encoded by nucleic acids that will hybridize to the sequences of the sequence listings, or their complements, as outlined herein.

In a preferred embodiment, when the CRC protein is to be used to generate antibodies, for example for immunotherapy, the CRC protein should share at least one epitope or determinant with the full length protein. By "epitope" or "determinant" herein is meant a portion of a protein which will generate and/or bind an antibody or T-cell receptor in the context of MHC. Thus, in most instances, antibodies made to a smaller CRC protein will be able to bind to the full length protein. In a preferred embodiment, the epitope is unique; that is, antibodies generated to a unique epitope show little or no cross-reactivity. In a preferred embodiment, the epitope is selected from CAA2p1 and CAA2p2. In another preferred embodiment, the epitope is selected from CAA3p1, CAA3p2, CAA3p3, CAAQ3p4, CAA3p4MAPS, CAA39p5 and CAA3p5MAPS.

In one embodiment, the term "antibody" includes antibody fragments, as are known in the art, including Fab, Fab₂, single chain antibodies (Fv for example), chimeric antibodies, etc., either produced by the modification of whole antibodies or those synthesized de novo using recombinant DNA technologies.

Methods of preparing polyclonal antibodies are known to the skilled artisan. Polyclonal antibodies can be raised in a mammal, for example, by one or more injections of an immunizing agent and, if desired, an adjuvant. Typically, the immunizing agent and/or adjuvant will be injected in the mammal by multiple subcutaneous or intraperitoneal injections. The immunizing agent may include the CAA2 or

fragment thereof or a fusion protein thereof. It may be useful to conjugate the immunizing agent to a protein known to be immunogenic in the mammal being immunized. Examples of such immunogenic proteins include but are not limited to keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, and soybean trypsin inhibitor. Examples of adjuvants which may be employed include Freund's complete adjuvant and MPL-TDM adjuvant (monophosphoryl Lipid A, synthetic trehalose dicorynomycolate). The immunization protocol may be selected by one skilled in the art without undue experimentation.

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The antibodies may, alternatively, be monoclonal antibodies. Monoclonal antibodies may be prepared using hybridoma methods, such as those described by Kohler and Milstein, Nature, 256:495 (1975). In a hybridoma method, a mouse, hamster, or other appropriate host animal, is typically immunized with an immunizing agent to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the immunizing agent. Alternatively, the lymphocytes may be immunized in vitro. The immunizing agent will typically include the CAA2 polypeptide or fragment thereof or a fusion protein thereof. Generally, either peripheral blood lymphocytes ("PBLs") are used if cells of human origin are desired, or spleen cells or lymph node cells are used if non-human mammalian sources are desired. The lymphocytes are then fused with an immortalized cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell [Goding, Monoclonal Antibodies: Principles and Practice, Academic Press, (1986) pp. 59-103]. Immortalized cell lines are usually transformed mammalian cells, particularly myeloma cells of rodent, bovine and human origin. Usually, rat or mouse myeloma cell lines are employed. The hybridoma cells may be cultured in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, immortalized cells. For example, if the parental cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine ("HAT medium"), which substances prevent the growth of HGPRT-deficient cells.

In one embodiment, the antibodies are bispecific antibodies. Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens. In the present case, one of the binding specificities is for a CRC protein or a fragment thereof, the other one is for any other antigen, and preferably for a cell-surface protein or receptor or receptor subunit, preferably one that is tumor specific.

In a preferred embodiment, the antibodies to CRC are capable of reducing or eliminating the biological function of CRC, as is described below. That is, the addition of anti-CRC antibodies (either polyclonal or preferably monoclonal) to CRC (or cells containing CRC) may reduce or eliminate the CRC activity.

Generally, at least a 25% decrease in activity is preferred, with at least about 50% being particularly preferred and about a 95-100% decrease being especially preferred.

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In a preferred embodiment the antibodies to the CRC proteins are humanized antibodies. Humanized forms of non-human (e.g., murine) antibodies are chimeric molecules of immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab'), or other antigen-binding subsequences of antibodies) which contain minimal sequence derived from non-human immunoglobulin. Humanized antibodies include human immunoglobulins (reciplent antibody) in which residues form a complementary determining region (CDR) of the recipient are replaced by residues from a CDR of a non-human species (donor antibody) such as mouse, rat or rabbit having the desired specificity, affinity and capacity. In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies may also comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the FR regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin [Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-329 (1988); and Presta, Curr. Op. Struct. Biol., 2:593-596 (1992)].

Methods for humanizing non-human antibodies are well known in the art. Generally, a humanized antibody has one or more amino acid residues introduced into it from a source which is non-human. These non-human amino acid residues are often referred to as import residues, which are typically taken from an import variable domain. Humanization can be essentially performed following the method of Winter and co-workers [Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-327 (1988); Verhoeyen et al., Science, 239:1534-1536 (1988)], by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such humanized antibodies are chimeric antibodies (U.S. Patent No. 4,816,567), wherein substantially less than an intact human variable domain has been substituted by the corresponding sequence from a non-human species. In practice, humanized antibodies are typically human antibodies in which some
 CDR residues and possibly some FR residues are substituted by residues from analogous sites in rodent antibodies.

Human antibodies can also be produced using various techniques known in the art, including phage display libraries [Hoogenboom and Winter, <u>J. Mol. Biol.</u>, 227:381 (1991); Marks et al., <u>J. Mol. Biol.</u>,

222:581 (1991)]. The techniques of Cole et al. and Boerner et al. are also available for the preparation of human monoclonal antibodies (Cole et al., Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, p. 77 (1985) and Boerner et al., J. Immunol., 147(1):86-95 (1991)]. Similarly, human antibodies can be made by introducing of human immunoglobulin loci into transgenic animals, e.g., mice in which the endogenous immunoglobulin genes have been partially or completely inactivated. Upon challenge, human antibody production is observed, which closely resembles that seen in humans in all respects, including gene rearrangement, assembly, and antibody repertoire. This approach is described, for example, in U.S. Patent Nos. 5,545,807; 5,545,806; 5,569,825; 5,625,126; 5,633,425; 5,661,016, and in the following scientific publications: Marks et al., Bio/Technology 10, 779-783 (1992); Lonberg et al., Nature 368, 856-859 (1994); Morrison, Nature 368, 812-13 (1994); Fishwild et al., Nature Biotechnology 14, 845-51 (1996); Neuberger, Nature Biotechnology 14, 826 (1996); Lonberg and Huszar, Intern. Rev. Immunol. 13 65-93 (1995).

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By immunotherapy is meant treatment of CRC with an antibody raised against CRC proteins. As used herein, immunotherapy can be passive or active. Passive immunotherapy as defined herein is the passive transfer of antibody to a recipient (patient). Active immunization is the induction of antibody and/or T-cell responses in a recipient (patient). Induction of an immune response is the result of providing the recipient with an antigen to which antibodies are raised. As appreciated by one of ordinary skill in the art, the antigen may be provided by injecting a polypeptide against which antibodies are desired to be raised into a recipient, or contacting the recipient with a nucleic acid capable of expressing the antigen and under conditions for expression of the antigen.

In a preferred embodiment the CRC proteins against which antibodies are raised are secreted proteins as described above. Without being bound by theory, antibodies used for treatment, bind and prevent the secreted protein from binding to its receptor, thereby inactivating the secreted CRC protein.

In another preferred embodiment, the CRC protein to which antibodies are raised is a transmembrane protein. Without being bound by theory, antibodies used for treatment, bind the extracellular domain of the CRC protein and prevent it from binding to other proteins, such as circulating ligands or cell-associated molecules. The antibody may cause down-regulation of the transmembrane CRC protein. As will be appreciated by one of ordinary skill in the art, the antibody may be a competitive, non-competitive or uncompetitive inhibitor of protein binding to the extracellular domain of the CRC protein. The antibody is also an antagonist of the CRC protein. Further, the antibody prevents activation of the transmembrane CRC protein. In one aspect, when the antibody prevents the binding of other molecules to the CRC protein, the antibody prevents growth of the cell. The antibody also sensitizes

the cell to cytotoxic agents, including, but not limited to TNF-a, TNF-b, IL-1, INF-g and IL-2, or chemotherapeutic agents including 5FU, vinblastine, actinomycin D, cisplatin, methotrexate, and the like. In some instances the antibody belongs to a sub-type that activates serum complement when complexed with the transmembrane protein thereby mediating cytotoxicity. Thus, CRC is treated by administering to a patient antibodies directed against the transmembrane CRC protein.

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In another preferred embodiment, the antibody is conjugated to a therapeutic moiety. In one aspect the therapeutic moiety is a small molecule that modulates the activity of the CRC protein. In another aspect the therapeutic moiety modulates the activity of molecules associated with or in close proximity to the CRC protein. The therapeutic moiety may inhibit enzymatic activity such as protease or protein kinase activity associated with CRC.

In a preferred embodiment, the therapeutic moiety may also be a cytotoxic agent. In this method, targeting the cytotoxic agent to tumor tissue or cells, results in a reduction in the number of afflicted cells, thereby reducing symptoms associated with CRC. Cytotoxic agents are numerous and varied and include, but are not limited to, cytotoxic drugs or toxins or active fragments of such toxins. Suitable toxins and their corresponding fragments include diptheria A chain, exotoxin A chain, ricin A chain, abrin A chain, curcin, crotin, phenomycin, enomycin and the like. Cytotoxic agents also include radiochemicals made by conjugating radioisotopes to antibodies raised against CRC proteins, or binding of a radionuclide to a chelating agent that has been covalently attached to the antibody. Targeting the therapeutic moiety to transmembrane CRC proteins not only serves to increase the local concentration of therapeutic moiety in the CRC afflicted area, but also serves to reduce deleterious side effects that may be associated with the therapeutic moiety.

In another preferred embodiment, the CRC protein against which the antibodies are raised is an intracellular protein. In this case, the antibody may be conjugated to a protein which facilitates entry into the cell. In one case, the antibody enters the cell by endocytosis. In another embodiment, a nucleic acid encoding the antibody is administered to the individual or cell. Moreover, wherein the CRC protein can be targeted within a cell, i.e., the nucleus, an antibody thereto contains a signal for that target localization, i.e., a nuclear localization signal.

The CRC antibodies of the Invention specifically bind to CRC proteins. By "specifically bind" herein is meant that the antibodies bind to the protein with a binding constant in the range of at least 10^{-4} - 10^{-8} M⁻¹, with a preferred range being 10^{-7} - 10^{-9} M⁻¹.

In a preferred embodiment, the CRC protein is purified or isolated after expression. CRC proteins may be isolated or purified in a variety of ways known to those skilled in the art depending on what other components are present in the sample. Standard purification methods include electrophoretic, molecular, immunological and chromatographic techniques, including ion exchange, hydrophobic, affinity, and reverse-phase HPLC chromatography, and chromatofocusing. For example, the CRC protein may be purified using a standard anti-CRC antibody column. Ultrafiltration and diafiltration techniques, in conjunction with protein concentration, are also useful. For general guidance in suitable purification techniques, see Scopes, R., Protein Purification, Springer-Verlag, NY (1982). The degree of purification necessary will vary depending on the use of the CRC protein. In some instances no purification will be necessary.

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Once expressed and purified if necessary, the CRC proteins and nucleic acids are useful in a number of applications.

In one aspect, the expression levels of genes are determined for different cellular states in the CRC phenotype; that is, the expression levels of genes in normal colon tissue and in CRC tissue (and in some cases, for varying severities of CRC that relate to prognosis, as outlined below) are evaluated to provide expression profiles. An expression profile of a particular cell state or point of development is essentially a "fingerprint" of the state; while two states may have any particular gene similarly expressed, the evaluation of a number of genes simultaneously allows the generation of a gene expression profile that is unique to the state of the cell. By comparing expression profiles of cells in different states, information regarding which genes are important (including both up- and down-regulation of genes) in each of these states is obtained. Then, diagnosis may be done or confirmed: does tissue from a particular patient have the gene expression profile of normal or CRC tissue.

"Differential expression," or grammatical equivalents as used herein, refers to both qualitative as well as quantitative differences in the genes' temporal and/or cellular expression patterns within and among the cells. Thus, a differentially expressed gene can qualitatively have its expression altered, including an activation or inactivation, in, for example, normal versus CRC tissue. That is, genes may be turned on or turned off in a particular state, relative to another state. As is apparent to the skilled artisan, any comparison of two or more states can be made. Such a qualitatively regulated gene will exhibit an expression pattern within a state or cell type which is detectable by standard techniques in one such state or cell type, but is not detectable in both. Alternatively, the determination is quantitative in that expression is increased or decreased; that is, the expression of the gene is either upregulated, resulting in an increased amount of transcript, or downregulated, resulting in a decreased amount of transcript. The degree to which expression differs need only be large enough to quantify via standard

characterization techniques as outlined below, such as by use of Affymetrix GeneChip™ expression arrays, Lockhart, Nature Biotechnology, 14:1675-1680 (1996), hereby expressly incorporated by reference. Other techniques include, but are not limited to, quantitative reverse transcriptase PCR, Northern analysis and RNase protection. As outlined above, preferably the change in expression (i.e. upregulation or downregulation) is at least about 50%, more preferably at least about 100%, more preferably at least about 150%, more preferably, at least about 200%, with from 300 to at least 1000% being especially preferred.

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As will be appreciated by those in the art, this may be done by evaluation at either the gene transcript, or the protein level; that is, the amount of gene expression may be monitored using nucleic acid probes to the DNA or RNA equivalent of the gene transcript, and the quantification of gene expression levels, or, alternatively, the final gene product itself (protein) can be monitored, for example through the use of antibodies to the CRC protein and standard immunoassays (ELISAs,e tc.) or other techniques, including mass spectroscopy assays, 2D gel electrophoresis assays, etc. Thus, the proteins corresponding to CRC genes, i.e. those identified as being important in a CRC phenotype, can be evaluated in a CRC diagnostic test.

In a preferred embodiment, gene expression monitoring is done and a number of genes, i.e. an expression profile, is monitored simultaneously, although multiple protein expression monitoring can be done as well. Similarly, these assays may be done on an individual basis as well.

In this embodiment, the CRC nucleic acid probes are attached to biochips as outlined herein for the detection and quantification of CRC sequences in a particular cell. The assays are further described below in the example.

In a preferred embodiment nucleic acids encoding the CRC protein are detected. Although DNA or RNA encoding the CRC protein may be detected, of particular interest are methods wherein the mRNA encoding a CRC protein is detected. The presence of mRNA in a sample is an indication that the CRC gene has been transcribed to form the mRNA, and suggests that the protein is expressed. Probes to detect the mRNA can be any nucleotide/deoxynucleotide probe that is complementary to and base pairs with the mRNA and includes but is not limited to oligonucleotides, cDNA or RNA. Probes also should contain a detectable label, as defined herein. In one method the mRNA is detected after immobilizing the nucleic acid to be examined on a solid support such as nylon membranes and hybridizing the probe with the sample. Following washing to remove the non-specifically bound probe, the label is detected. In another method detection of the mRNA is performed in situ. In this method permeabilized cells or tissue samples are contacted with a detectably labeled

nucleic acid probe for sufficient time to allow the probe to hybridize with the target mRNA. Following washing to remove the non-specifically bound probe, the label is detected. For example a digoxygenin labeled riboprobe (RNA probe) that is complementary to the mRNA encoding a CRC protein is detected by binding the digoxygenin with an anti-digoxygenin secondary antibody and developed with nitro blue tetrazolium and 5-bromo-4-chloro-3-indoyl phosphate.

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In a preferred embodiment, any of the three classes of proteins as described herein (secreted, transmembrane or intracellular proteins) are used in diagnostic assays. The CRC proteins, antibodies, nucleic acids, modified proteins and cells containing CRC sequences are used in diagnostic assays. This can be done on an individual gene or corresponding polypeptide level. In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes and/or corresponding polypeptides.

As described and defined herein, CRC proteins, including intracellular, transmembrane or secreted proteins, find use as markers of CRC. Detection of these proteins in putative CRC tissue or patients allows for a determination or diagnosis of CRC. Numerous methods known to those of ordinary skill in the art find use in detecting CRC. In one embodiment, antibodies are used to detect CRC proteins. A preferred method separates proteins from a sample or patient by electrophoresis on a gel (typically a denaturing and reducing protein gel, but may be any other type of gel including isoelectric focusing gels and the like). Following separation of proteins, the CRC protein is detected by immunoblotting with antibodies raised against the CRC protein. Methods of immunoblotting are well known to those of ordinary skill in the art.

In another preferred method, antibodies to the CRC protein find use in in situ imaging techniques. In this method cells are contacted with from one to many antibodies to the CRC protein(s). Following washing to remove non-specific antibody binding, the presence of the antibody or antibodies is detected. In one embodiment the antibody is detected by incubating with a secondary antibody that contains a detectable label. In another method the primary antibody to the CRC protein(s) contains a detectable label. In another preferred embodiment each one of multiple primary antibodies contains a distinct and detectable label. This method finds particular use in simultaneous screening for a plurality of CRC proteins. As will be appreciated by one of ordinary skill in the art, numerous other histological imaging techniques are useful in the invention.

In a preferred embodiment the label is detected in a fluorometer which has the ability to detect and distinguish emissions of different wavelengths. In addition, a fluorescence activated cell sorter (FACS) can be used in the method.

In another preferred embodiment, antibodies find use in diagnosing CRC from blood samples. As previously described, certain CRC proteins are secreted/circulating molecules. Blood samples, therefore, are useful as samples to be probed or tested for the presence of secreted CRC proteins. Antibodies can be used to detect the CRC by any of the previously described immunoassay techniques including ELISA, immunoblotting (Western blotting), immunoprecipitation, BIACORE technology and the like, as will be appreciated by one of ordinary skill in the art.

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In a preferred embodiment, in situ hybridization of labeled CRC nucleic acid probes to tissue arrays is done. For example, arrays of tissue samples, including CRC tissue and/or normal tissue, are made.

In situ hybridization as is known in the art can then be done.

It is understood that when comparing the fingerprints between an individual and a standard, the skilled artisan can make a diagnosis as well as a prognosis. It is further understood that the genes which indicate the diagnosis may differ from those which indicate the prognosis.

In a preferred embodiment, the CRC proteins, antibodies, nucleic acids, modified proteins and cells containing CRC sequences are used in prognosis assays. As above, gene expression profiles can be generated that correlate to CRC severity, in terms of long term prognosis. Again, this may be done on either a protein or gene level, with the use of genes being preferred. As above, the CRC probes are attached to biochips for the detection and quantification of CRC sequences in a tissue or patient. The assays proceed as outlined for diagnosis.

In a preferred embodiment, any of the three classes of proteins as described herein are used in drug screening assays. The CRC proteins, antibodies, nucleic acids, modified proteins and cells containing CRC sequences are used in drug screening assays or by evaluating the effect of drug candidates on a "gene expression profile" or expression profile of polypeptides. In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes after treatment with a candidate agent, Zlokamik, et al., Science 279, 84-8 (1998), Heid, 1996 #69.

In a preferred embodiment, the CRC proteins, antibodies, nucleic acids, modified proteins and cells containing the native or modified CRC proteins are used in screening assays. That is, the present

invention provides novel methods for screening for compositions which modulate the CRC phenotype. As above, this can be done on an individual gene level or by evaluating the effect of drug candidates on a "gene expression profile". In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes after treatment with a candidate agent, see Zlokarnik, supra.

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Having identified the differentially expressed genes herein, a variety of assays may be executed. In a preferred embodiment, assays may be run on an individual gene or protein level. That is, having identified a particular gene as up regulated in CRC, candidate bioactive agents may be screened to modulate this gene's response; preferably to down regulate the gene, although in some circumstances to up regulate the gene. "Modulation" thus includes both an increase and a decrease in gene expression. The preferred amount of modulation will depend on the original change of the gene expression in normal versus tumor tissue, with changes of at least 10%, preferably 50%, more preferably 100-300%, and in some embodiments 300-1000% or greater. Thus, if a gene exhibits a 4 fold increase in tumor compared to normal tissue, a decrease of about four fold is desired; a 10 fold decrease in tumor compared to normal tissue gives a 10 fold increase in expression for a candidate agent is desired.

As will be appreciated by those in the art, this may be done by evaluation at either the gene or the protein level; that is, the amount of gene expression may be monitored using nucleic acid probes and the quantification of gene expression levels, or, alternatively, the gene product itself can be monitored, for example through the use of antibodies to the CRC protein and standard immunoassays.

In a preferred embodiment, gene expression monitoring is done and a number of genes, i.e. an expression profile, is monitored simultaneously, although multiple protein expression monitoring can be done as well.

In this embodiment, the CRC nucleic acid probes are attached to biochips as outlined herein for the detection and quantification of CRC sequences in a particular cell. The assays are further described below.

Generally, in a preferred embodiment, a candidate bioactive agent is added to the cells prior to analysis. Moreover, screens are provided to identify a candidate bioactive agent which modulates colorectal cancer, modulates CRC proteins, binds to a CRC protein, or interferes between the binding of a CRC protein and an antibody.

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The term "candidate bioactive agent" or "drug candidate" or grammatical equivalents as used herein describes any molecule, e.g., protein, oligopeptide, small organic molecule, polysaccharide, polynucleotide, etc., to be tested for bioactive agents that are capable of directly or indirectly altering either the CRC phenotype or the expression of a CRC sequence, including both nucleic acid sequences and protein sequences. In preferred embodiments, the bioactive agents modulate the expression profiles, or expression profile nucleic acids or proteins provided herein. In a particularly preferred embodiment, the candidate agent suppresses a CRC phenotype, for example to a normal colon tissue fingerprint. Similarly, the candidate agent preferably suppresses a severe CRC phenotype. Generally a plurality of assay mixtures are run in parallel with different agent concentrations to obtain a differential response to the various concentrations. Typically, one of these concentrations serves as a negative control, i.e., at zero concentration or below the level of detection.

In one aspect, a candidate agent will neutralize the effect of a CRC protein. By "neutralize" is meant that activity of a protein is either inhibited or counter acted against so as to have substantially no effect on a cell

Candidate agents encompass numerous chemical classes, though typically they are organic molecules, preferably small organic compounds having a molecular weight of more than 100 and less than about 2,500 daltons. Preferred small molecules are less than 2000, or less than 1500 or less than 1000 or less than 500 D. Candidate agents comprise functional groups necessary for structural interaction with proteins, particularly hydrogen bonding, and typically include at least an amine, carbonyl, hydroxyl or carboxyl group, preferably at least two of the functional chemical groups. The candidate agents often comprise cyclical carbon or heterocyclic structures and/or aromatic or polyaromatic structures substituted with one or more of the above functional groups. Candidate agents are also found among biomolecules including peptides, saccharides, fatty acids, steroids, purines, pyrimidines, derivatives, structural analogs or combinations thereof. Particularly preferred are peptides.

Candidate agents are obtained from a wide variety of sources including libraries of synthetic or natural compounds. For example, numerous means are available for random and directed synthesis of a wide variety of organic compounds and biomolecules, including expression of randomized oligonucleotides. Alternatively, libraries of natural compounds in the form of bacterial, fungal, plant and animal extracts are available or readily produced. Additionally, natural or synthetically produced libraries and compounds are readily modified through conventional chemical, physical and biochemical means. Known pharmacological agents may be subjected to directed or random

chemical modifications, such as acylation, alkylation, esterification, amidification to produce structural analogs.

In a preferred embodiment, the candidate bioactive agents are proteins. By "protein" herein is meant at least two covalently attached amino acids, which includes proteins, polypeptides, oligopeptides and peptides. The protein may be made up of naturally occurring amino acids and peptide bonds, or synthetic peptidomimetic structures. Thus "amino acid", or "peptide residue", as used herein means both naturally occurring and synthetic amino acids. For example, homo-phenylalanine, citrulline and noreleucine are considered amino acids for the purposes of the invention. "Amino acid" also includes imino acid residues such as proline and hydroxyproline. The side chains may be in either the (R) or the (S) configuration. In the preferred embodiment, the amino acids are in the (S) or L-configuration. If non-naturally occurring side chains are used, non-amino acid substituents may be used, for example to prevent or retard in vivo degradations.

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In a preferred embodiment, the candidate bioactive agents are naturally occurring proteins or fragments of naturally occurring proteins. Thus, for example, cellular extracts containing proteins, or random or directed digests of proteinaceous cellular extracts, may be used. In this way libraries of procaryotic and eucaryotic proteins may be made for screening in the methods of the invention. Particularly preferred in this embodiment are libraries of bacterial, fungal, viral, and mammalian proteins, with the latter being preferred, and human proteins being especially preferred.

In a preferred embodiment, the candidate bioactive agents are peptides of from about 5 to about 30 amino acids, with from about 5 to about 20 amino acids being preferred, and from about 7 to about 15 being particularly preferred. The peptides may be digests of naturally occurring proteins as is outlined above, random peptides, or "biased" random peptides. By "randomized" or grammatical equivalents herein is meant that each nucleic acid and peptide consists of essentially random nucleotides and amino acids, respectively. Since generally these random peptides (or nucleic acids, discussed below) are chemically synthesized, they may incorporate any nucleotide or amino acid at any position. The synthetic process can be designed to generate randomized proteins or nucleic acids, to allow the formation of all or most of the possible combinations over the length of the sequence, thus forming a library of randomized candidate bioactive proteinaceous agents.

In one embodiment, the library is fully randomized, with no sequence preferences or constants at any position. In a preferred embodiment, the library is biased. That is, some positions within the sequence are either held constant, or are selected from a limited number of possibilities. For example, in a preferred embodiment, the nucleotides or amino acid residues are randomized within a

defined class, for example, of hydrophobic amino acids, hydrophilic residues, sterically biased (either small or large) residues, towards the creation of nucleic acid binding domains, the creation of cysteines, for cross-linking, prolines for SH-3 domains, serines, threonines, tyrosines or histidines for phosphorylation sites, etc., or to purines, etc.

In a preferred embodiment, the candidate bioactive agents are nucleic acids, as defined above.

As described above generally for proteins, nucleic acid candidate bioactive agents may be naturally occurring nucleic acids, random nucleic acids, or "biased" random nucleic acids. For example, digests of procaryotic or eucaryotic genomes may be used as is outlined above for proteins.

In a preferred embodiment, the candidate bioactive agents are organic chemical moieties, a wide variety of which are available in the literature.

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After the candidate agent has been added and the cells allowed to incubate for some period of time, the sample containing the target sequences to be analyzed is added to the biochip. If required, the target sequence is prepared using known techniques. For example, the sample may be treated to lyse the cells, using known lysis buffers, electroporation, etc., with purification and/or amplification such as PCR occurring as needed, as will be appreciated by those in the art. For example, an in vitro transcription with labels covalently attached to the nucleosides is done. Generally, the nucleic acids are labeled with biotin-FITC or PE, or with cy3 or cy5.

In a preferred embodiment, the target sequence is labeled with, for example, a fluorescent, a chemiluminescent, a chemical, or a radioactive signal, to provide a means of detecting the target sequence's specific binding to a probe. The label also can be an enzyme, such as, alkaline phosphatase or horseradish peroxidase, which when provided with an appropriate substrate produces a product that can be detected. Alternatively, the label can be a labeled compound or small molecule, such as an enzyme inhibitor, that binds but is not catalyzed or altered by the enzyme. The label also can be a molety or compound, such as, an epitope tag or biotin which specifically binds to streptavidin. For the example of biotin, the streptavidin is labeled as described above, thereby, providing a detectable signal for the bound target sequence. As known in the art, unbound labeled streptavidin is removed prior to analysis.

As will be appreciated by those in the art, these assays can be direct hybridization assays or can comprise "sandwich assays", which include the use of multiple probes, as is generally outlined in U.S. Patent Nos. 5,681,702, 5,597,909, 5,545,730, 5,594,117, 5,591,584, 5,571,670, 5,580,731, 5,571,670,

5,591,584, 5,624,802, 5,635,352, 5,594,118, 5,359,100, 5,124,246 and 5,681,697, all of which are hereby incorporated by reference. In this embodiment, in general, the target nucleic acid is prepared as outlined above, and then added to the biochip comprising a plurality of nucleic acid probes, under conditions that allow the formation of a hybridization complex.

A variety of hybridization conditions may be used in the present invention, including high, moderate and low stringency conditions as outlined above. The assays are generally run under stringency conditions which allows formation of the label probe hybridization complex only in the presence of target. Stringency can be controlled by altering a step parameter that is a thermodynamic variable, including, but not limited to, temperature, formamide concentration, salt concentration, chaotropic salt concentration pH, organic solvent concentration, etc.

These parameters may also be used to control non-specific binding, as is generally outlined in U.S. Patent No. 5,681,697. Thus it may be desirable to perform certain steps at higher stringency conditions to reduce non-specific binding.

The reactions outlined herein may be accomplished in a variety of ways, as will be appreciated by those in the art. Components of the reaction may be added simultaneously, or sequentially, in any order, with preferred embodiments outlined below. In addition, the reaction may include a variety of other reagents may be included in the assays. These include reagents like salts, buffers, neutral proteins, e.g. albumin, detergents, etc which may be used to facilitate optimal hybridization and detection, and/or reduce non-specific or background interactions. Also reagents that otherwise improve the efficiency of the assay, such as protease inhibitors, nuclease inhibitors, anti-microbial agents, etc., may be used, depending on the sample preparation methods and purity of the target.

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Once the assay is run, the data is analyzed to determine the expression levels, and changes in expression levels as between states, of individual genes, forming a gene expression profile.

The screens are done to identify drugs or bioactive agents that modulate the CRC phenotype. Specifically, there are several types of screens that can be run. A preferred embodiment is in the screening of candidate agents that can induce or suppress a particular expression profile, thus preferably generating the associated phenotype. That is, candidate agents that can mimic or produce an expression profile in CRC similar to the expression profile of normal colon tissue is expected to result in a suppression of the CRC phenotype. Thus, in this embodiment, mimicking an expression profile, or changing one profile to another, is the goal.

In a preferred embodiment, as for the diagnosis and prognosis applications, having identified the differentially expressed genes important in any one state, screens can be run to alter the expression of the genes individually. That is, screening for modulation of regulation of expression of a single gene can be done; that is, rather than try to mimic all or part of an expression profile, screening for regulation of individual genes can be done. Thus, for example, particularly in the case of target genes whose presence or absence is unique between two states, screening is done for modulators of the target gene expression.

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In a preferred embodiment, screening is done to alter the biological function of the expression product of the differentially expressed gene. Again, having identified the importance of a gene in a particular state, screening for agents that bind and/or modulate the biological activity of the gene product can be run as is more fully outlined below.

Thus, screening of candidate agents that modulate the CRC phenotype either at the gene expression level or the protein level can be done.

In addition screens can be done for novel genes that are induced in response to a candidate agent. After identifying a candidate agent based upon its ability to suppress a CRC expression pattern leading to a normal expression pattern, or modulate a single CRC gene expression profile so as to mimic the expression of the gene from normal tissue, a screen as described above can be performed to identify genes that are specifically modulated in response to the agent. Comparing expression profiles between normal tissue and agent treated CRC tissue reveals genes that are not expressed in normal tissue or CRC tissue, but are expressed in agent treated tissue. These agent specific sequences can be identified and used by any of the methods described herein for CRC genes or proteins. In particular these sequences and the proteins they encode find use in marking or identifying agent treated cells. In addition, antibodies can be raised against the agent induced proteins and used to target novel therapeutics to the treated CRC tissue sample.

Thus, in one embodiment, a candidate agent is administered to a population of CRC cells, that thus has an associated CRC expression profile. By "administration" or "contacting" herein is meant that the candidate agent is added to the cells in such a manner as to allow the agent to act upon the cell, whether by uptake and intracellular action, or by action at the cell surface. In some embodiments, nucleic acid encoding a proteinaceous candidate agent (i.e. a peptide) may be put into a viral construct such as a retroviral construct and added to the cell, such that expression of the peptide agent is accomplished; see PCT US97/01019, hereby expressly incorporated by reference.

Once the candidate agent has been administered to the cells, the cells can be washed if desired and are allowed to incubate under preferably physiological conditions for some period of time. The cells are then harvested and a new gene expression profile is generated, as outlined herein.

Thus, for example, CRC tissue may be screened for agents that reduce or suppress the CRC phenotype. A change in at least one gene of the expression profile indicates that the agent has an effect on CRC activity. By defining such a signature for the CRC phenotype, screens for new drugs that alter the phenotype can be devised. With this approach, the drug target need not be known and need not be represented in the original expression screening platform, nor does the level of transcript for the target protein need to change.

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In a preferred embodiment, as outlined above, screens may be done on individual genes and gene products (proteins). That is, having identified a particular differentially expressed gene as important in a particular state, screening of modulators of either the expression of the gene or the gene product itself can be done. The gene products of differentially expressed genes are sometimes referred to herein as "CRC proteins" or a "CCMP". In preferred embodiments, the CCMP is as depicted in Figures 17-20, 24, 25, 29, 33 and 36, more preferably the protein having the sequence shown in Figures 29 or 36 or encoded by the sequences of Figures 27, 36 and 39-48. The CCMP may be a fragment, or alternatively, be the full length protein to a fragment shown herein. Preferably, the CCMP is a fragment of approximately 14 to 24 amino acids long. More preferably the fragment is a soluble fragment.

In a preferred embodiment, the fragment is from CAA9. Preferably, the fragment includes a non-transmenbrane region. In a preferred embodiment, the CAA9 fragment has an N-terminal Cys to aid in solubility. Preferably, the fragment is selected from CAA9p1, Caa9p2, CAA9p3, CAA9p4, CAA9p4MAPS, CAA9p5 and CAA9p5MAPS.

In a preferred embodiment, the fragment is charged and from the c-terminus of CAA2. In one embodiment, the c-terminus of the fragment is kept as a free acid and the n-terminus is a free amine to aid in coupling, i.e., to cysteine. In another embodiment, the fragment is an internal peptide overlapping hydrophilic stretch of CAA2. In a preferred embodiment, the termini is blocked. Preferably, the fragment of CAA2 is selected from CAA2p1 or CAA2p2. In another preferred embodiment, the fragment is a novel fragment from the N-terminal. In one embodiment, the fragment excludes sequence outside of the N-terminal, in another embodiment, the fragment includes at least a portion of the N-terminal. "N-terminal" is used interchangeably herein with "N-terminus" which is further described above.

In one embodiment the CRC proteins are conjugated to an immunogenic agent as discussed herein. In one embodiment the CRC protein is conjugated to BSA.

Thus, in a preferred embodiment, screening for modulators of expression of specific genes can be done. This will be done as outlined above, but in general the expression of only one or a few genes are evaluated.

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In a preferred embodiment, screens are designed to first find candidate agents that can bind to differentially expressed proteins, and then these agents may be used in assays that evaluate the ability of the candidate agent to modulate differentially expressed activity. Thus, as will be appreciated by those in the art, there are a number of different assays which may be run; binding assays and activity assays.

In a preferred embodiment, binding assays are done. In general, purified or isolated gene product is used; that is, the gene products of one or more differentially expressed nucleic acids are made. In general, this is done as is known in the art. For example, antibodies are generated to the protein gene products, and standard immunoassays are run to determine the amount of protein present.

Alternatively, cells comprising the CRC proteins can be used in the assays.

Thus, in a preferred embodiment, the methods comprise combining a CRC protein and a candidate bioactive agent, and determining the binding of the candidate agent to the CRC protein. Preferred embodiments utilize the human CRC protein, although other mammalian proteins may also be used, for example for the development of animal models of human disease. In some embodiments, as outlined herein, variant or derivative CRC proteins may be used.

Generally, in a preferred embodiment of the methods herein, the CRC protein or the candidate agent is non-diffusably bound to an insoluble support having isolated sample receiving areas (e.g. a microtiter plate, an array, etc.). The insoluble supports may be made of any composition to which the compositions can be bound, is readily separated from soluble material, and is otherwise compatible with the overall method of screening. The surface of such supports may be solid or porous and of any convenient shape. Examples of suitable insoluble supports include microtiter plates, arrays, membranes and beads. These are typically made of glass, plastic (e.g., polystyrene), polysaccharides, nylon or nitrocellulose, teflonTM, etc. Microtiter plates and arrays are especially convenient because a large number of assays can be carried out simultaneously, using small amounts of reagents and samples. The particular manner of binding of the composition is not crucial so long as it is compatible with the reagents and overall methods of the invention, maintains the activity of the

composition and is nondiffusable. Preferred methods of binding include the use of antibodies (which do not sterically block either the ligand binding site or activation sequence when the protein is bound to the support), direct binding to "sticky" or ionic supports, chemical crosslinking, the synthesis of the protein or agent on the surface, etc. Following binding of the protein or agent, excess unbound material is removed by washing. The sample receiving areas may then be blocked through incubation with bovine serum albumin (BSA), casein or other innocuous protein or other moiety.

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In a preferred embodiment, the CRC protein is bound to the support, and a candidate bioactive agent is added to the assay. Alternatively, the candidate agent is bound to the support and the CRC protein is added. Novel binding agents include specific antibodies, non-natural binding agents identified in screens of chemical libraries, peptide analogs, etc. Of particular interest are screening assays for agents that have a low toxicity for human cells. A wide variety of assays may be used for this purpose, including labeled in vitro protein-protein binding assays, electrophoretic mobility shift assays, immunoassays for protein binding, functional assays (phosphorylation assays, etc.) and the like.

The determination of the binding of the candidate bioactive agent to the CRC protein may be done in a number of ways. In a preferred embodiment, the candidate bioactive agent is labeled, and binding determined directly. For example, this may be done by attaching all or a portion of the CRC protein to a solid support, adding a labeled candidate agent (for example a fluorescent label), washing off excess reagent, and determining whether the label is present on the solid support. Various blocking and washing steps may be utilized as is known in the art.

By "labeled" herein is meant that the compound is either directly or indirectly labeled with a label which provides a detectable signal, e.g. radioisotope, fluorescers, enzyme, antibodies, particles such as magnetic particles, chemiluminescers, or specific binding molecules, etc. Specific binding molecules include pairs, such as biotin and streptavidin, digoxin and antidigoxin etc. For the specific binding members, the complementary member would normally be labeled with a molecule which provides for detection, in accordance with known procedures, as outlined above. The label can directly or indirectly provide a detectable signal.

In some embodiments, only one of the components is labeled. For example, the proteins (or proteinaceous candidate agents) may be labeled at tyrosine positions using ¹²⁵I, or with fluorophores. Alternatively, more than one component may be labeled with different labels; using ¹²⁵I for the proteins, for example, and a fluorophor for the candidate agents.

In a preferred embodiment, the binding of the candidate bioactive agent is determined through the use of competitive binding assays. In this embodiment, the competitor is a binding moiety known to bind to the target molecule (i.e. CRC), such as an antibody, peptide, binding partner, ligand, etc. Under certain circumstances, there may be competitive binding as between the bioactive agent and the binding moiety, with the binding moiety displacing the bioactive agent.

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In one embodiment, the candidate bloactive agent is labeled. Either the candidate bloactive agent, or the competitor, or both, is added first to the protein for a time sufficient to allow binding, if present. Incubations may be performed at any temperature which facilitates optimal activity, typically between 4 and 40°C. Incubation periods are selected for optimum activity, but may also be optimized to facilitate rapid high through put screening. Typically between 0.1 and 1 hour will be sufficient. Excess reagent is generally removed or washed away. The second component is then added, and the presence or absence of the labeled component is followed, to indicate binding.

In a preferred embodiment, the competitor is added first, followed by the candidate bioactive agent. Displacement of the competitor is an indication that the candidate bioactive agent is binding to the CRC protein and thus is capable of binding to, and potentially modulating, the activity of the CRC protein. In this embodiment, either component can be labeled. Thus, for example, if the competitor is labeled, the presence of label in the wash solution indicates displacement by the agent. Alternatively, if the candidate bloactive agent is labeled, the presence of the label on the support indicates displacement.

In an alternative embodiment, the candidate bioactive agent is added first, with incubation and washing, followed by the competitor. The absence of binding by the competitor may indicate that the bioactive agent is bound to the CRC protein with a higher affinity. Thus, if the candidate bioactive agent is labeled, the presence of the label on the support, coupled with a lack of competitor binding, may indicate that the candidate agent is capable of binding to the CRC protein.

In a preferred embodiment, the methods comprise differential screening to identity bloactive agents that are capable of modulating the activity of the CRC proteins. In this embodiment, the methods comprise combining a CRC protein and a competitor in a first sample. A second sample comprises a candidate bloactive agent, a CRC protein and a competitor. The binding of the competitor is determined for both samples, and a change, or difference in binding between the two samples indicates the presence of an agent capable of binding to the CRC protein and potentially modulating its activity. That is, if the binding of the competitor is different in the second sample relative to the first sample, the agent is capable of binding to the CRC protein.

Alternatively, a preferred embodiment utilizes differential screening to identify drug candidates that bind to the native CRC protein, but cannot bind to modified CRC proteins. The structure of the CRC protein may be modeled, and used in rational drug design to synthesize agents that interact with that site. Drug candidates that affect CRC bioactivity are also identified by screening drugs for the ability to either enhance or reduce the activity of the protein.

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Positive controls and negative controls may be used in the assays. Preferably all control and test samples are performed in at least triplicate to obtain statistically significant results. Incubation of all samples is for a time sufficient for the binding of the agent to the protein. Following incubation, all samples are washed free of non-specifically bound material and the amount of bound, generally labeled agent determined. For example, where a radiolabel is employed, the samples may be counted in a scintillation counter to determine the amount of bound compound.

A variety of other reagents may be included in the screening assays. These include reagents like salts, neutral proteins, e.g. albumin, detergents, etc which may be used to facilitate optimal protein-protein binding and/or reduce non-specific or background interactions. Also reagents that otherwise improve the efficiency of the assay, such as protease inhibitors, nuclease inhibitors, anti-microbial agents, etc., may be used. The mixture of components may be added in any order that provides for the requisite binding.

Screening for agents that modulate the activity of CRC proteins may also be done. In a preferred embodiment, methods for screening for a bioactive agent capable of modulating the activity of CRC proteins comprise the steps of adding a candidate bioactive agent to a sample of CRC proteins, as above, and determining an alteration in the biological activity of CRC proteins. "Modulating the activity of CRC" includes an increase in activity, a decrease in activity, or a change in the type or kind of activity present. Thus, in this embodiment, the candidate agent should both bind to CRC proteins (although this may not be necessary), and alter its biological or biochemical activity as defined herein. The methods include both in vitro screening methods, as are generally outlined above, and in vivo screening of cells for alterations in the presence, distribution, activity or amount of CRC proteins.

Thus, in this embodiment, the methods comprise combining a CRC sample and a candidate bioactive agent, and evaluating the effect on CRC activity. By "CRC activity" or grammatical equivalents herein is meant one of the CRC's biological activities, including, but not limited to, cell division, preferably in colon tissue, cell proliferation, tumor growth, transformation of cells. In one embodiment, CRC activity includes activation of CZA8, BCX2, CBC2, CBC1, CBC3, CJA9, BCN5, CQA1, BCN7, CQA2, CJA8,

CAA2, CAA9, CGA7 and/or CGA8*, preferably one of the CRC proteins listed in Figure 14. An inhibitor of CRC activity is the inhibition of any one or more CRC activities.

In a preferred embodiment, the activity of the CRC protein is increased; in another preferred embodiment, the activity of the CRC protein is decreased. Thus, bioactive agents that are antagonists are preferred in some embodiments, and bioactive agents that are agonists may be preferred in other embodiments.

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In a preferred embodiment, the invention provides methods for screening for bioactive agents capable of modulating the activity of a CRC protein. The methods comprise adding a candidate bioactive agent, as defined above, to a cell comprising CRC proteins. Preferred cell types include almost any cell. The cells contain a recombinant nucleic acid that encodes a CRC protein. In a preferred embodiment, a library of candidate agents are tested on a plurality of cells.

In one aspect, the assays are evaluated in the presence or absence or previous or subsequent exposure of physiological signals, for example hormones, antibodies, peptides, antigens, cytokines, growth factors, action potentials, pharmacological agents including chemotherapeutics, radiation, carcinogenics, or other cells (i.e. cell-cell contacts). In another example, the determinations are determined at different stages of the cell cycle process.

In this way, bioactive agents are identified. Compounds with pharmacological activity are able to enhance or interfere with the activity of the CRC protein. In one embodiment, "colorectal cancer protein activity" as used herein includes at least one of the following: colorectal cancer activity, binding to CJA8, activation of CJA8 or activation of substrates of CJA8 by CJA8. In one embodiment, colorectal cancer activity is defined as the unregulated proliferation of colon tissue, or the growth of cancer in colon tissue. In one aspect, colorectal cancer activity as defined herein is related to the activity of CJA8 in the upregulation of CJA8 in colon cancer tissue.

In another embodiment, colorectal cancer protein activity includes at least one of the following: colorectal cancer activity, binding to one of CAA2, CAA9, CGA7 and CGA8, activation of one of CAA2, CAA9, CGA7, and CGA8 or activation of substrates of CAA2, CAA9, CGA7 or CGA8 by CAA2, CAA9, CGA7 or CGA8, respectively. In one preferred embodiment, CAA2 comprises its N-terminal end. In one aspect, colorectal cancer activity as defined herein is related to the activity of CAA2, CAA9, CGA7 and/or CGA8 in the upregulation of CAA2, CAA9, CGA7 and/or CGA8, respectively, in colon cancer tissue.

In one embodiment, a method of inhibiting colon cancer cell division is provided. The method comprises administration of a colorectal cancer inhibitor.

In another embodiment, a method of inhibiting tumor growth is provided. The method comprises administration of a colorectal cancer inhibitor.

In a further embodiment, methods of treating cells or individuals with cancer are provided. The method comprises administration of a colorectal cancer inhibitor.

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In one embodiment, a colorectal cancer inhibitor is an antibody as discussed above. In another embodiment, the colorectal cancer inhibitor is an antisense molecule. Antisense molecules as used herein include antisense or sense oligonucleotides comprising a singe-stranded nucleic acid sequence (either RNA or DNA) capable of binding to target mRNA (sense) or DNA (antisense) sequences for colorectal cancer molecules. A preferred antisense molecule is for CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, BCN5, CQA1, BCN7, CQA2, CAA2, CAA9, CGA7 or CGA8, more preferably for the CRC sequences referenced in Figure 14, or for a ligand or activator thereof. A most preferred antisense molecule is for CJA8 or for a ligand or activator thereof. Antisense or sense oligonucleotides, according to the present invention, comprise a fragment generally at least about 14 nucleotides, preferably from about 14 to 30 nucleotides. The ability to derive an antisense or a sense oligonucleotide, based upon a cDNA sequence encoding a given protein is described in, for example, Stein and Cohen (Cancer Res. 48:2659, 1988) and van der Krol et al. (BioTechniques 6:958, 1988).

Antisense molecules may be introduced into a cell containing the target nucleotide sequence by formation of a conjugate with a ligand binding molecule, as described in WO 91/04753. Suitable ligand binding molecules include, but are not limited to, cell surface receptors, growth factors, other cytokines, or other ligands that bind to cell surface receptors. Preferably, conjugation of the ligand binding molecule does not substantially interfere with the ability of the ligand binding molecule to bind to its corresponding molecule or receptor, or block entry of the sense or antisense oligonucleotide or its conjugated version into the cell. Alternatively, a sense or an antisense oligonucleotide may be introduced into a cell containing the target nucleic acid sequence by formation of an oligonucleotide-lipid complex, as described in WO 90/10448. It is understood that the use of antisense molecules or knock out and knock in models may also be used in screening assays as discussed above, in addition to methods of treatment.

The compounds having the desired pharmacological activity may be administered in a physiologically acceptable carrier to a host, as previously described. The agents may be administered in a variety of

ways, orally, parenterally e.g., subcutaneously, intraperitoneally, intravascularly, etc. Depending upon the manner of introduction, the compounds may be formulated in a variety of ways. The concentration of therapeutically active compound in the formulation may vary from about 0.1-100 wt.%. The agents may be administered alone or in combination with other treatments, i.e., radiation.

The pharmaceutical compositions can be prepared in various forms, such as granules, tablets, pills, suppositories, capsules, suspensions, salves, lotions and the like. Pharmaceutical grade organic or inorganic carriers and/or diluents suitable for oral and topical use can be used to make up compositions containing the therapeutically-active compounds. Diluents known to the art include aqueous media, vegetable and animal oils and fats. Stabilizing agents, wetting and emulsifying agents, salts for varying the osmotic pressure or buffers for securing an adequate pH value, and skin penetration enhancers can be used as auxiliary agents.

Without being bound by theory, it appears that the various CRC sequences are important in CRC. Accordingly, disorders based on mutant or variant CRC genes may be determined. In one embodiment, the invention provides methods for identifying cells containing variant CRC genes comprising determining all or part of the sequence of at least one endogeneous CRC genes in a cell. As will be appreciated by those in the art, this may be done using any number of sequencing techniques. In a preferred embodiment, the invention provides methods of identifying the CRC genotype of an individual comprising determining all or part of the sequence of at least one CRC gene of the individual. This is generally done in at least one tissue of the individual, and may include the evaluation of a number of tissues or different samples of the same tissue. The method may include comparing the sequence of the sequenced CRC gene to a known CRC gene, i.e. a wild-type gene.

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The sequence of all or part of the CRC gene can then be compared to the sequence of a known CRC gene to determine if any differences exist. This can be done using any number of known homology programs, such as Bestfit, etc. In a preferred embodiment, the presence of a a difference in the sequence between the CRC gene of the patient and the known CRC gene is indicative of a disease state or a propensity for a disease state, as outlined herein.

In a preferred embodiment, the CRC genes are used as probes to determine the number of copies of the CRC gene in the genome.

In another preferred embodiment CRC genes are used as probed to determine the chromosomal localization of the CRC genes. Information such as chromosomal localization finds use in providing a

diagnosis or prognosis in particular when chromosomal abnormalities such as translocations, and the like are identified in CRC gene loci.

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Thus, in one embodiment, methods of modulating CRC in cells or organisms are provided. In one embodiment, the methods comprise administering to a cell an anti-CRC antibody that reduces or eliminates the biological activity of an endogeneous CRC protein. Alternatively, the methods comprise administering to a cell or organism a recombinant nucleic acid encoding a CRC protein. As will be appreciated by those in the art, this may be accomplished in any number of ways. In a preferred embodiment, for example when the CRC sequence is down-regulated in CRC, the activity of the CRC gene is increased by increasing the amount of CRC in the cell, for example by overexpressing the endogeneous CRC or by administering a gene encoding the CRC sequence, using known genetherapy techniques, for example. In a preferred embodiment, the gene therapy techniques include the incorporation of the erogenous gene using enhanced homologous recombination (EHR), for example as described in PCT/US93/03868, hereby incorporated by reference in its entirety. Alternatively, for example when the CRC sequence is up-regulated in CRC, the activity of the endogeneous CRC gene is decreased, for example by the administration of a CRC antisense nucleic acid.

In one embodiment, the CRC proteins of the present invention may be used to generate polyclonal and monoclonal antibodies to CRC proteins, which are useful as described herein. Similarly, the CRC proteins can be coupled, using standard technology, to affinity chromatography columns. These columns may then be used to purify CRC antibodies. In a preferred embodiment, the antibodies are generated to epitopes unique to a CRC protein; that is, the antibodies show little or no cross-reactivity to other proteins. These antibodies find use in a number of applications. For example, the CRC antibodies may be coupled to standard affinity chromatography columns and used to purify CRC proteins. The antibodies may also be used as blocking polypeptides, as outlined above, since they will specifically bind to the CRC protein.

In one embodiment, a therapeutically effective dose of a CRC or modulator thereof is administered to a patient. By "therapeutically effective dose" herein is meant a dose that produces the effects for which it is administered. The exact dose will depend on the purpose of the treatment, and will be ascertainable by one skilled in the art using known techniques. As is known in the art, adjustments for CRC degradation, systemic versus localized delivery, and rate of new protease synthesis, as well as the age, body weight, general health, sex, diet, time of administration, drug interaction and the severity of the condition may be necessary, and will be ascertainable with routine experimentation by those skilled in the art.

A "patient" for the purposes of the present invention includes both humans and other animals, particularly mammals, and organisms. Thus the methods are applicable to both human therapy and veterinary applications. In the preferred embodiment the patient is a mammal, and in the most preferred embodiment the patient is human.

The administration of the CRC proteins and modulators of the present invention can be done in a variety of ways as discussed above, including, but not limited to, orally, subcutaneously, intravenously, intranasally, transdermally, intraperitoneally, intramuscularly, intrapulmonary, vaginally, rectally, or intraocularly. In some instances, for example, in the treatment of wounds and inflammation, the CRC proteins and modulators may be directly applied as a solution or spray.

10 The pharmaceutical compositions of the present invention comprise a CRC protein in a form suitable for administration to a patient. In the preferred embodiment, the pharmaceutical compositions are in a water soluble form, such as being present as pharmaceutically acceptable salts, which is meant to include both acid and base addition salts. "Pharmaceutically acceptable acid addition salt" refers to those salts that retain the biological effectiveness of the free bases and that are not biologically or otherwise undesirable, formed with inorganic acids such as hydrochloric acid, hydrobromic acid, 15 sulfuric acid, nitric acid, phosphoric acid and the like, and organic acids such as acetic acid, propionic acid, glycolic acid, pyruvic acid, oxalic acid, maleic acid, malonic acid, succinic acid, fumaric acid, tartaric acid, citric acid, benzoic acid, cinnamic acid, mandelic acid, methanesulfonic acid, ethanesulfonic acid, p-toluenesulfonic acid, salicylic acid and the like. "Pharmaceutically acceptable 20 base addition salts" include those derived from inorganic bases such as sodium, potassium, lithium, ammonium, calcium, magnesium, iron, zinc, copper, manganese, aluminum salts and the like. Particularly preferred are the ammonium, potassium, sodium, calcium, and magnesium salts. Salts derived from pharmaceutically acceptable organic non-toxic bases include salts of primary, secondary, and tertiary amines, substituted amines including naturally occurring substituted amines, cyclic amines 25 and basic ion exchange resins, such as isopropylamine, trimethylamine, diethylamine, triethylamine, tripropylamine, and ethanolamine.

The pharmaceutical compositions may also include one or more of the following: carrier proteins such as serum albumin; buffers; fillers such as microcrystalline cellulose, lactose, corn and other starches; binding agents; sweeteners and other flavoring agents; coloring agents; and polyethylene glycol. Additives are well known in the art, and are used in a variety of formulations.

In a preferred embodiment, CRC proteins and modulators are administered as therapeutic agents, and can be formulated as outlined above. Similarly, CRC genes (including both the full-length

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sequence, partial sequences, or regulatory sequences of the CRC coding regions) can be administered in gene therapy applications, as is known in the art. These CRC genes can include antisense applications, either as gene therapy (i.e. for incorporation into the genome) or as antisense compositions, as will be appreciated by those in the art.

In a preferred embodiment, CRC genes are administered as DNA vaccines, either single genes or combinations of CRC genes. Naked DNA vaccines are generally known in the art. Brower, Nature Biotechnology, 16:1304-1305 (1998).

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In one embodiment, CRC genes of the present invention are used as DNA vaccines. Methods for the use of genes as DNA vaccines are well known to one of ordinary skill in the art, and include placing a CRC gene or portion of a CRC gene under the control of a promoter for expression in a CRC patient. The CRC gene used for DNA vaccines can encode full-length CRC proteins, but more preferably encodes portions of the CRC proteins including peptides derived from the CRC protein. In a preferred embodiment a patient is immunized with a DNA vaccine comprising a plurality of nucleotide sequences derived from a CRC gene. Similarly, it is possible to immunize a patient with a plurality of CRC genes or portions thereof as defined herein. Without being bound by theory, expression of the polypeptide encoded by the DNA vaccine, cytotoxic T-cells, helper T-cells and antibodies are induced which recognize and destroy or eliminate cells expressing CRC proteins.

In a preferred embodiment, the DNA vaccines include a gene encoding an adjuvant molecule with the DNA vaccine. Such adjuvant molecules include cytokines that increase the immunogenic response to the CRC polypeptide encoded by the DNA vaccine. Additional or alternative adjuvants are known to those of ordinary skill in the art and find use in the invention.

In another preferred embodiment CRC genes find use in generating animal models of CRC. As is appreciated by one of ordinary skill in the art, when the CRC gene identified is repressed or diminished in CRC tissue, gene therapy technology wherein antisense RNA directed to the CRC gene will also diminish or repress expression of the gene. An animal generated as such serves as an animal model of CRC that finds use in screening bioactive drug candidates. Similarly, gene knockout technology, for example as a result of homologous recombination with an appropriate gene targeting vector, will result in the absence of the CRC protein. When desired, tissue-specific expression or knockout of the CRC protein may be necessary.

It is also possible that the CRC protein is overexpressed in CRC. As such, transgenic animals can be generated that overexpress the CRC protein. Depending on the desired expression level, promoters

of various strengths can be employed to express the transgene. Also, the number of copies of the integrated transgene can be determined and compared for a determination of the expression level of the transgene. Animals generated by such methods find use as animal models of CRC and are additionally useful in screening for bioactive molecules to treat CRC.

It is understood that the examples described herein in no way serve to limit the true scope of this invention, but rather are presented for illustrative purposes. All references and sequences of accession numbers cited herein are incorporated by reference in their entirety.

EXAMPLES

Example 1

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10 <u>Tissue Preparation, Labeling Chips, and Fingerprints</u>

Purify total RNA from tissue using TRIzol Reagent

Estimate tissue weight. Homogenize tissue samples in 1ml of TRIzol per 50mg of tissue using a Polytron 3100 homogenizer. The generator/probe used depends upon the tissue size. A generator that is too large for the amount of tissue to be homogenized will cause a loss of sample and lower RNA yield. Use the 20mm generator for tissue weighing more than 0.6g. If the working volume is greater than 2ml, then homogenize tissue in a 15ml polypropylene tube (Falcon 2059). Fill tube no greater than 10ml.

HOMOGENIZATION

Before using generator, it should have been cleaned after last usage by running it through soapy H20 and rinsing thoroughly. Run through with EtOH to sterilize. Keep tissue frozen until ready. Add TRIzol directly to frozen tissue then homogenize.

Following homogenization, remove insoluble material from the homogenate by centrifugation at 7500 x g for 15 min. in a Sorvall superspeed or 12,000 x g for 10 min. in an Eppendorf centrifuge at 4°C. Transfer the cleared homogenate to a new tube(s). The samples may be frozen now at -60 to -70°C (and kept for at least one month) or you may continue with the purification.

PHASE SEPARATION

Incubate the homogenized samples for 5 minutes at room temperature.

Add 0.2ml of chloroform per 1ml of TRIzol reagent used in the original homogenization. Cap tubes securely and shake tubes vigorously by hand (do not vortex) for 15 seconds.

Incubate samples at room temp. for 2-3 minutes. Centrifuge samples at $^{\prime}$ 5500rpm in a Sorvall superspeed for 30 min. at 4°C. (You may spin at up to 12,000 x g for 10 min. but you risk breaking your tubes in the centrifuge.)

RNA PRECIPITATION

Transfer the aqueous phase to a fresh tube. Save the organic phase if isolation of DNA or protein is desired. Add 0.5ml of isopropyl alcohol per 1ml of TRIzol reagent used in the original homogenization. Cap tubes securely and invert to mix. Incubate samples at room temp. for 10 minutes. Centrifuge samples at 6500rpm in Sorvall for 20min. at 4°C.

RNA WASH

Pour off the supernate. Wash pellet with cold 75% ethanol. Use 1ml of 75% ethanol per 1ml of TRIzol reagent used in the initial homogenization. Cap tubes securely and invert several times to loosen pellet. (Do not vortex). Centrifuge at <8000rpm (<7500 x g) for 5 minutes at 4°C. Pour off the wash. Carefully transfer pellet to an eppendorf tube (let it slide down the tube into the new tube and use a pipet tip to help guide it in if necessary). Depending on the volumes you are working with, you can decide what size tube(s) you want to precipitate the RNA in. When I tried leaving the RNA in the large 15ml tube, it took so long to dry (i.e. it did not dry) that I eventually had to transfer it to a smaller tube. Let pellet dry in hood. Resuspend RNA in an appropriate volume of DEPC H₂0. Try for 2-5ug/ul. Take absorbance readings.

Purify poly A+ mRNA from total RNA or clean up total RNA with Qiagen's

20 RNeasy kit

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Purification of poly A* mRNA from total RNA. Heat oligotex suspension to 37°C and mix immediately before adding to RNA. Incubate Elution Buffer at 70°C. Warm up 2 x Binding Buffer at 65°C if there is precipitate in the buffer. Mix total RNA with DEPC-treated water, 2 x Binding Buffer, and Oligotex according to Table 2 on page 16 of the Oligotex Handbook. Incubate for 3 minutes at 65°C. Incubate for 10 minutes at room temperature.

Centrifuge for 2 minutes at 14,000 to 18,000 g. If centrifuge has a "soft setting," then use it. Remove supernatant without disturbing Oligotex pellet. A little bit of solution can be left behind to reduce the loss of Oligotex. Save sup until certain that satisfactory binding and elution of poly A* mRNA has occurred.

Gently resuspend in Wash Buffer OW2 and pipet onto spin column. Centrifuge the spin column at full speed (soft setting if possible) for 1 minute.

Transfer spin column to a new collection tube and gently resuspend in Wash Buffer OW2 and centrifuge as describe herein.

Transfer spin column to a new tube and elute with 20 to 100 ul of preheated (70°C) Elution Buffer. Gently resuspend Oligotex resin by pipetting up and down. Centrifuge as above. Repeat elution with fresh elution buffer or use first eluate to keep the elution volume low.

Read absorbance, using diluted Elution Buffer as the blank.

Before proceeding with cDNA synthesis, the mRNA must be precipitated.

Some component leftover or in the Elution Buffer from the Oligotex purification procedure will inhibit downstream enzymatic reactions of the mRNA.

10 Ethanol Precipitation

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Add 0.4 vol. of 7.5 M NH₄OAc + 2.5 vol. of cold 100% ethanol. Precipitate at -20°C 1 hour to overnight (or 20-30 min. at -70°C). Centrifuge at 14,000-16,000 x g for 30 minutes at 4°C. Wash pellet with 0.5ml of 80%ethanol (-20°C) then centrifuge at 14,000-16,000 x g for 5 minutes at room temperature. Repeat 80% ethanol wash. Dry the last bit of ethanol from the pellet in the hood. (Do not speed vacuum). Suspend pellet in DEPC H₂0 at 1ug/ul concentration.

Clean up total RNA using Qiagen's RNeasy kit

Add no more than 100ug to an RNeasy column. Adjust sample to a volume of 100ul with RNase-free water. Add 350ul Buffer RLT then 250ul ethanol (100%) to the sample. Mix by pipetting (do not centrifuge) then apply sample to an RNeasy mini spin column. Centrifuge for 15 sec at >10,000rpm. If concerned about yield, re-apply flowthrough to column and centrifuge again. Transfer column to a new 2-ml collection tube. Add 500ul Buffer RPE and centrifuge for 15 sec at >10,000rpm. Discard flowthrough. Add 500ul Buffer RPE and centrifuge for 15 sec at >10,000rpm. Discard flowthrough then centrifuge for 2 min at maximum speed to dry column membrane. Transfer column to a new 1.5-ml collection tube and apply 30-50ul of RNase-free water directly onto column membrane. Centrifuge 1 min at >10,000rpm. Repeat elution. Take absorbance reading. If necessary, ethanol precipitate with ammonium acetate and 2.5X volume 100% ethanol.

Make cDNA using Gibco's "SuperScript Choice System for cDNA Synthesis" kit First Strand cDNA Synthesis

Use 5ug of total RNA or 1ug of polyA+ mRNA as starting material. For total RNA, use 2ul of SuperScript RT. For polyA+ mRNA, use 1ul of SuperScript RT. Final volume of first strand synthesis mix is 20ul. RNA must be in a volume no greater than 10ul. Incubate RNA with 1ul of 100pmol T7-T24 oligo for 10 min at 70C. On ice, add 7 ul of: 4ul 5X 1st Strand Buffer, 2ul of 0.1M DTT, and 1 ul of 10mM dNTP mix. Incubate at 37C for 2 min then add SuperScript RT Incubate at 37C for 1 hour.

Second Strand Synthesis

Place 1st strand reactions on ice.

Add:

91ul DEPC H20

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30ul 5X 2nd Strand Buffer

3ul 10mM dNTP mix

1ul 10U/ul E.coli DNA Ligase

4ul 10U/ul E.coli DNA Polymerase

1ul 2U/ul RNase H

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Make the above into a mix if there are more than 2 samples. Mix and incubate 2 hours at 16C. Add 2ul T4 DNA Polymerase. Incubate 5 min at 16C. Add 10ul of 0.5M EDTA

Clean up cDNA

Phenol:Chloroform:Isoamyl Alcohol (25:24:1) purification using Phase-Lock gel tubes:

Centrifuge PLG tubes for 30 sec at maximum speed. Transfer cDNA mix to PLG tube. Add equal volume of phenol:chloroform:isamyl alcohol and shake vigorously (do not vortex). Centrifuge 5 minutes at maximum speed. Transfer top aqueous solution to a new tube. Ethanol precipitate: add 7.5X 5M NH4Oac and 2.5X volume of 100% ethanol. Centrifuge immediately at room temp. for 20 min, maximum speed. Remove sup then wash pellet 2X with cold 80% ethanol. Remove as much ethanol wash as possible then let pellet air dry. Resuspend pellet in 3ul RNase-free water.

In vitro Transcription (IVT) and labeling with biotin

Pipet 1.5ul of cDNA into a thin-wall PCR tube.

Make NTP labeling mix:

30 Combine at room temperature: 2ul

T7 10xATP (75mM) (Ambion)

2ul

T7 10xGTP (75mM) (Ambion)

1.5ul

T7 10xCTP (75mM) (Ambion)

1.5ul T7 10xUTP (75mM) (Ambion)

3.75ul 10mM Bio-11-UTP (Boehringer-Mannheim/Roche or

Enzo)

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3.75ul 10mM Bio-16-CTP (Enzo) -

2ul 10x T7 transcription buffer (Ambion)

2ul 10x T7 enzyme mix (Ambion)

Final volume of total reaction is 20ul. Incubate 6 hours at 37C in a PCR machine.

RNeasy clean-up of IVT product

Follow previous instructions for RNeasy columns or refer to Qiagen's RNeasy protocol handbook.

cRNA will most likely need to be ethanol precipitated. Resuspend in a volume compatible with the fragmentation step.

Fragmentation

15 ug of labeled RNA is usually fragmented. Try to minimize the fragmentation reaction volume;
a 10 ul volume is recommended but 20 ul is all right. Do not go higher than 20 ul because the
magnesium in the fragmentation buffer contributes to precipitation in the hybridization buffer.
Fragment RNA by incubation at 94 C for 35 minutes in 1 x Fragmentation buffer.

5 x Fragmentation buffer:

200 mM Tris-acetate, pH 8.1 500 mM KOAc

20 150 mM MgOAc

The labeled RNA transcript can be analyzed before and after fragmentation. Samples can be heated to 65C for 15 minutes and electrophoresed on 1% agarose/TBE gels to get an approximate idea of the transcript size range

Hybridization

25 200 ul (10ug cRNA) of a hybridization mix is put on the chip. If multiple hybridizations are to be done (such as cycling through a 5 chip set), then it is recommended that an initial hybridization mix of 300 ul or more be made. Hybrization Mix: fragment labeled RNA (50ng/ul final conc.)

50 pM 948-b control oligo

1.5 pM BioB

5 pM BioC

25 pM BioD

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100 pM CRE

0.1mg/ml herring sperm DNA

0.5mg/ml acetylated BSA

to 300 ul with 1xMES hyb. buffer

The instruction manuals for the products used herein are incorporated herein in their entirety.

Labeling Protocol Provided Herein

Hybridization reaction:

Start with non-biotinylated IVT (purified by RNeasy columns)

(see example 1 for steps from tissue to IVT)

15 IVT antisense RNA; 4 μg:

ρ.

Random Hexamers (1 µg/µl): 4 µl

H₂O:

μi

14 µI

- Incubate 70°C, 10 min. Put on ice.

Reverse transcription:

5X First Strand (BRL) buffer: 6 µl

0.1 M DTT:

3 µ1

50X dNTP mix:

0.6 µl

25 H2O:

2.4 µl

Cy3 or Cy5 dUTP (1mM):

3 µl

SS RT II (BRL):

1 µl

16 µl

30 - Add to hybridization reaction.

- Incubate 30 min., 42°C.

- Add 1 µl SSII and let go for another hour.

Put on ice.

- 50X dNTP mix (25mM of cold dATP, dCTP, and dGTP, 10mM of dTTP: 25 µl each of 100mM dATP, dCTP, and dGTP; 10 µl of 100mM dTTP to 15 µl H2O. dNTPs from Pharmacia)

RNA degradation:

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86 µl H₂O

10 μl 10N NaOH

- Add 1.5 µl 1M NaOH/ 2mM EDTA, incubate at 65°C, 10 min.

4 µl 50mM EDTA

U-Con 30

500 µl TE/sample spin at 7000g for 10 min, save flow through for purification

10 Qiagen purification:

- -suspend u-con recovered material in 500µl buffer PB
- -proceed w/ normal Qiagen protocol

DNAse digest:

- Add 1 µl of 1/100 dil of DNAse/30µl Rx and incubate at 37°C for 15 min.
- 15 -5 min 95°C to denature enzyme

Sample preparation:

- Add:

Cot-1 DNA: 10 µI

50X dNTPs: 1 µl

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20X SSC: 2.3 µl

Na pyro phosphate: 7.5 µl 10mg/ml Herring sperm DNA 1ul of 1/10 dilution

21.8 final vol.

- Dry down in speed vac.
- 25 Resuspend in 15 μl H₂0.
 - Add 0.38 µl 10% SDS.
 - Heat 95°C, 2 min.
 - Slow cool at room temp. for 20 min.

Put on slide and hybridize overnight at 64°C.

30 Washing after the hybridization:

3X SSC/0.03% SDS:

2 min. 37.5 mls 20X SSC+0.75mls 10% SDS in 250mls H₂O

1X SSC: 5 min.

12.5 mls 20X SSC in 250mls H_2O

0.2X SSC: 5 min.

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2.5 mls 20X SSC in 250mls H2O

Dry slides in centrifuge, 1000 RPM, 1min. Scan at appropriate PMT's and channels.

The results are shown in Figures 1 through 11. The lists of genes come from colorectal tumors from a variety of stages of the disease. The genes that are up regulated in the tumors (overall) were also found to be expressed at a limited amount or not at all in the body map. The body map for the colorectal project consists of ten tissues: Heart, Brain, Lung, Liver, Breast, Kidney, Prostrate, Small Intestine, Spleen, and Colon. The down regulated genes in tumors (overall) versus normal colon were not selected for their expression or lack of expression in the body map. As indicated, some of the Accession numbers include expression sequence tags (ESTs). Thus, in one embodiment herein, genes within an expression profile, also termed expression profile genes, include ESTs and are not necessarily full length. Figure 1 shows 51 upregulated genes; Figure 2 shows 194 upregulated genes; Figure 3 shows 1144 upregulated genes and Figure 4 shows 1815 upregulated genes. The genes shown in Figures 1 and 5 are particularly preferred. Figure 5 shows 54 downregulated genes; Figure 6 shows 558 downregulated genes; and Figure 7 shows 1923 downregulated genes; and Figures 8, 9, 10 and 11 provide the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal colon tissue.

Example 2

20 Expression studies were performed herein.

As indicated in Figure 21, CAA2 is upregulated in colon cancer tissue. CAA2 is found in chromosome 15, cytoband 15q15-22, interval D15S146-D15S117. CAA2 has N-myristoylation sites and a C-terminal microbody targeting signal. The preferred fragments shown in Figures 18 and 19 have a solubility of 1 mg/ 1 ml H20.

As indicated in Figure 26, CAA9 is upregulated in colon cancer tissue. CAA9 is found in chromosome 5, cytoband 5q23.3, interval D5S471-D5S393.

As indicated in Figures 30A and 30B, CGA7 is upregulated in colon cancer tissue. CGA7 is found in chromosome 2.

As indicated in Figure 34, CGA8 is upregulated in colon cancer tissue.

As indicated in Figure 37, CJA8 is upregulated in colon cancer tissue. CJA8 is found in chromosome 11. $\,\cdot\,$

As indicated in Figure 38, BCN7 is upregulated in colon cancer tissue. BCN7 is found in chromosome 5, cytoband 5q22, interval D5S471-D5S393.

CLAIMS

We claim:

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- A method of screening drug candidates comprising:
 - a) providing a cell that expresses an expression profile gene which encodes a protein selected from the group consisting of CZA8, BCX2, CBC2, CBC1, CBC3, CJA8, CJA9, CGA7, BCN5, CQA1, BCN7 and CQA2 or a fragment thereof;
 - b) adding a drug candidate to said cell; and
 - c) determining the effect of said drug candidate on the expression of said expression profile gene.
- 2. A method according to claim 1 wherein said determining comprises comparing the level of expression in the absence of said drug candidate to the level of expression in the presence of said drug candidate, wherein the concentration of said drug candidate can vary when present, and wherein said comparison can occur after addition or removal of the drug candidate.
 - 3. A method according to claim 1 wherein the expression of said profile gene is decreased as a result of the introduction of the drug candidate.
 - 4. A method of screening for a bioactive agent capable of binding to a colorectal cancer modulator protein (CCMP), wherein said CCMP is CJA8 or a fragment thereof, said method comprising combining said CCMP and a candidate bioactive agent, and determining the binding of said candidate agent to said CCMP.
- 5. A method for screening for a bioactive agent capable of modulating the activity of a colorectal cancer modulator protein (CCMP), wherein said CCMP is CJA8 or a fragment thereof, said method comprising combining said CCMP and a candidate bioactive agent, and determining the effect of said candidate agent on the bioactivity of said CCMP.
 - 6. A method of evaluating the effect of a candidate colorectal cancer drug comprising:
- a) administering said drug to a patient;
 - b) removing a cell sample from said patient; and
 - c) determining the expression profile of said cell.
 - 7. A method according to claim 6 further comprising comparing said expression profile to an expression profile of a healthy individual.

8. A biochip comprising a nucleic acid segment encoding CJA81 or a fragment thereof, wherein said biochip comprises fewer than 1000 nucleic acid probes.

9. A method of diagnosing colorectal cancer comprising:

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- a) determining the expression of a gene encoding CJA8 or a fragment thereof in a first tissue type of a first individual; and
- b) comparing said expression of said gene from a second normal tissue type from said first individual or a second unaffected individual;

wherein a difference in said expression indicates that the first individual has colorectal cancer.

- 10. An antibody which specifically binds to CJA8, or a fragment thereof.
- 10 11. An antibody which specifically binds to CAA9, or a fragment thereof.
 - 12. The antibody of Claim 11 wherein said fragment is selected from the group CAA9p1, CAA9p2, CAA9p3, CAA9p4, CAA9p4MAPS, CAA9p5 and CAA9p5MAPS.
 - 13. The antibody of Claim 10, wherein said antibody is a monoclonal antibody.
 - 14. The antibody of Claim 10, wherein said antibody is a humanized antibody.
- 15. The antibody of Claim 10, wherein said antibody is an antibody fragment.
 - 16. A method for screening for a bioactive agent capable of interfering with the binding of a colorectal cancer modulator protein (CCMP) or a fragment thereof and an antibody which binds to said CCMP or fragment thereof, said method comprising:
 - a) combining a CCMP or fragment thereof, a candidate bioactive agent and an antibody which binds to said CCMP or fragment thereof; and
 - b) determining the binding of said CCMP or fragment thereof and said antibody.
 - 17. A method for inhibiting colorectal cancer, said method comprising administering to a cell a composition comprising an antibody to CAJ8 or a fragment thereof.
 - 18. The method of Claim 17 wherein said cell is a cell of an individual.
- 25 19. The method of Claim 18 wherein said individual has cancer.

20. The method of Claim 17 wherein said antibody is a humanized antibody.

- 21. The method of Claim 17 wherein said antibody is an antibody fragment.
- 22. A method for inhibiting colorectal cancer in a cell, wherein said method comprises administering to a cell a composition comprising antisense molecules to CJA8.
- 5 23. A peptide consisting essentially of CAA9p1, CAA9p2, CAA9p3, CAA9p4, CAA9p4MAPS, CAA9p5 or CAA9p5MAPS.
 - 24. A composition comprising the peptide of Claim 23.
 - 25. A method of eliciting an immune response in an individual, said method comprising administering to said individual a composition comprising CJA8 or a fragment thereof.
- 26. A method of eliciting an immune response in an individual, said method comprising administering to said individual a composition comprising a nucleic acid comprising a sequence encoding CJA8 or a fragment thereof.
 - 27. A composition capable of eliciting an immune response in an individual, said composition comprising CJA8 or a fragment thereof and a pharmaceutically acceptable carrier.
- 28. A composition capable of eliciting an immune response in an individual, said composition comprising a nucleic acid comprising a sequence encoding CJA8 or a fragment thereof and a pharmaceutically acceptable carrier.
 - 29. A method of treating an individual for colorectal cancer comprising administering to said individual an inhibitor of CJA8.
- 20 30. The method of Claim 29 wherein said inhibitor is an antibody.
 - 31. A method for determining the prognosis of an individual with colorectal cancer comprising determining the level of CJA8 in a sample, wherein a high level of CJA8 indicates a poor prognosis.

32. A method of neutralizing the effect of a CJA8, or a fragment thereof, comprising contacting an agent specific for said protein with said protein in an amount sufficient to effect neutralization.

- 33. A method for localizing a therapeutic molety to colorectal cancer tissue comprising exposing said tissue to an antibody to CJA8 or fragment thereof conjugated to said therapeutic molety.
 - 34. The method of Claim 33, wherein said therapeutic moiety is a cytotoxic agent.

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- 35. The method of Claim 33, wherein said therapeutic moiety is a radioisotope.
- 36. A method of treating colorectal cancer comprising administering to an individual having
 colorectal cancer an antibody to CJA8 or fragment thereof conjugated to a therapeutic moiety.
 - 37. The method of Claim 36, wherein said therapeutic moiety is a cytotoxic agent.
 - 38. The method of Claim 36, wherein said therapeutic molety is a radioisotope.

FIGURE 1

in Tumor over normal colon > 10	Accession AA460530	Unigene CLUSTER Hs.98384	Unigere Descriptor Homo sapiens orphan G protein-coupled receptor HG38 mRNA complete cds
	783366 U30246	Hs. 110736	n.sapiens micha for NBK apoptotic inducer protein Human burnetanide-sensitive Na-K-Cl cotransporter (NKCC1) mRNA complete cds
	AA216722	Hs.54481	Human mRNA for apolipoprotein E receptor 2 complete cds
	X17644	Hs.2707	G1 to S phase transition 1
	AA053636	Hs.129849	PBK1
	W59961	Hs.22564	Human mRNA for KIAA0389 gene complete cds
	AA449311	Hs.98658	Homo sapiens mitotic checkpoint protein kinase (BUB1) mRNA complete cds
	MS8597	Hs.2173	Fucosyltransferase 4 (alpha (13) fucosyltransferase myeloid-specific)
	AA598712	Hs.23723	ESTs Weakly similar to ORF YPL212c [S.cerevisiae]
	U55206	Hs.78619	Homo sapiens human gamma-glutamyl hydrolase (hGH) mRNA complete cds
	L48211	Hs.20954	Homo Sapiens angiotensIn II receptor gene complete cds
	X54942	Hs.83758	CDC28 protein kinase 2
	AA199747	Hs.79025	Human mRNA for KIAA0096 gene partial cds
	X68314	Hs.2704	Glutathione peroxidase 2 gastrointestinal
	AA405098	Hs.38178	ESTs Weakly similar to MOESIN/EZRIN/RADIXIN HOMOLOG (D.melanogaster)
	AA417030	Hs.5101	Homo saptens protein regulating cytokinesis 1 (PRC1) mRNA complete cds
	HG2981-HT3127		EST - HG2981-HT3127
	L41939	Hs.89403	Homo sapiens protein-tyrosine kinase EPHB2v (EPHB2) mRNA complete cds
	AA442763	Hs.20483	ESTs Highly similar to G2/MITOTIC-SPECIFIC CYCLIN B2 [Mesocricetus auratus]
	X17620	Hs.118638	NUCLEOSIDE DIPHOSPHATE KINASE A
	AA479139	Hs.75393	Acid phosphatase 1 soluble
	U48807	Hs.2359	Human MAP kinase phosphatase (MKP-2) mRNA complete cds
	AA243133	Hs.48915	Homo sapiens serine/threonine kinase (BTAK) mRNA complete cds
	H10984	Hs.12338	ESTs
	X54489	Hs. 789	GRO1 oncogene (melanoma growth stimulating activity alpha)
	W93943	Hs.59509	ESTs
	U48705	Hs.75562	Receptor protein-tyrosine kinase EDDR1
	X62048	Hs.75188	WEE1-LIKE PROTEIN KINASE
	AA505133	Hs.62273	ESTs
	R09195	Hs.151385	Hs.151385 Homo saplens mRNA for KIAA0564 protein partial cds

Hs.41723 Human kinesin-like spindle protein HKSP (HKSP) mRNA complete cds	· ESTs	EST - M30448	Human tumor necrosis factor type 1 receptor associated protein (TRAP1) mRNA padial cds			ESTs Weakly similar to renin [H.sapiens]			Transketolase (Wernicke-Korsakoff syndrome)										
Hs.41723	Hs,110457 ESTs		Hs.2204	Hs.93121	Hs.89690	Hs.25863	Hs.75761	Hs.42650	Hs.89643	Hs.8130	Hs.80706	Hs.7756	Hs.41697	Hs.79305	Hs.79217	Hs.80896	Hs.2236	Hs.79748	
AA453159	AA046745	M30448	U12595	U54999	X53800	AA011134	T29681	AA292765	L12711	AA113149	J03934	AA203428	D79997	D87444	M77836	Z29066	Z29067	M21904	
5.2	2.5	2.4	2.3	2.3	2.3	2.3	2.3	2.3	2.1	2.1	1.8	1.5	1.5	1.5	1.2	T	1.0	0.7	,
71148	25310	2640	3834	4674	5769	25050	41935	26895	1782	25593	1487	7656	683	836	3098	6819	6880	2473	00000

FIGURE 2

Unigene Descriptor	Professe inhibitor 5 (maspin)	Homo sapiens clone 23797 and 23917 mRNA partial cds	w	Human protein-tyrosine phosphatase (HU-PP-1) mRNA partial sequence	Ribonuckease L. (2'5'-oligoisoadenylate synthetase-dependent) inhibitor		w	Human mRNA for transcriptional activator hSNF2b complete cds	w	w	TONE H2A.X	Human DNA polymerase delta small subunit mRNA complete cds	Human bumetanide-sensitive Na-K-CI cotransporter (NKCC1) mRNA complete cds	Protease inhibitor 5 (maspin)		EST - HG4716-HT5158	ESTs Highly similar to CHROMOSOME CONDENSATION PROTEIN DPY-27 (Caenorhabditis elegans)	Homo sapiens mRNA for high mobility group protein HMG2a	Homo sapiens human gamma-glulamyl hydrolase (hGH) mRNA complete cds		N.	CDC28 protein kinase 2	, w	, w	FARNESYL-DIPHOSPHATE FARNESYLTRANSFERASE	Msh (Drosophila) homeo box homidog 2	- J05614	ESTs Highly similar to 60S RtBOSOMAL PROTEIN L22 [Rattus norvegicus]	· o	Glutathione peroxidase 2 gastrointestinal	ESTs Weakly similar (c. 3LU SUBFAMILY J) : [H.sapiens]	wlus]	Human antioxidant enzyme AOE37-2 mRNA complete cds		
Inigene Descri	rotease inhibit	forma sapiens	ESTs	fuman protein-	libonuclease L	EST	ESTs	furnan mRNA	ESTs	ESTs	HISTONE H2A.X	fuman DNA po	fuman bumeta	rotease inhibi	ESTs	ST - HG4716	STs Highly s	lomo sapiens	tomo sapiens	EST	ESTs	DC28 protein	ESTs	ESTs	ARNESYL-DI	Ash (Drosophil	EST - J05614	STs Highly s	ESTs	Slutathione per	STs Weakly	STs Moderate	fuman antioxic	ESTs	FST - 1191327
Unigene CLUSTER	Hs.55279 F	Hs.12540 F	Hs.31656 E	Hs.82911 F	Hs.12013 F	Hs.108240 E	Hs.22858 E	Hs.78202 F	Hs.101074 E	Hs.71190 E	Hs.2711 P	Hs.74598 F	Hs.110736 P	Hs.55279 F	Hs.47378 E	w	Hs.87630 E	Hs.19114 H	Hs.78619 F	Hs.25276 E	Hs.34782 E	_	Hs.124215 E	Hs.112013 E	Hs.48876 F	Hs.89404 N	.	Hs.54602 E	Hs.18457 E	Hs.2704 (Hs.19322 E	Hs.73625 E	Hs.83383 F	Hs.86430 E	u
Accession	W93726	AA232315	H20128	AA330771	AA223912	H62474	H17808	AA598648	AA504343	AA128407	X14850	U21090	U30246	U04313	AA331393	HG4716-HT5158	AA455239	AA435840	US5206	R38239	D59894	X54942	N22107	H03686	X69141	D89377	J05614	AA129757	N22015	X68314	AA088458	AA179845	U25182	AA211901	1101327
fold upregulated of Tumor	ر د د	۲۱0 ۲۷	v 10	V10	10	×10	<u>۲</u> 10	710	>10	>10	> 10	>10	۷. د ا	۷1۷	>10	۲٠ ۲0	10.0	9.6	9.6	8.7	83	4.7	7.1	7.0	7.0	8.8	6.7	6.7	6.4	6.1	6.1	0.9	5.9	5.8	e u
Primary Key	33616	34197	19387	8125	18362	39995	19328	38590	38456	17559	5619	4029	15006	3659	26916	1346	37491	13110	4676	21655	14723	5793	29848	9347	6078	9326	1566	25675	20126	6061	10867	18062	4093	18290	400

6928	5.7	746620	He 2316	SDV from deforministics and the state of the
6160	5.6	X74794	Hs.89699	CDC21 HOMOLOG
31487	54	N69507	Hs.129849	ESTS
9470	5.3	H46617		EST - H46617
33458	5.0	W86835	Hs.14158	Homo sapiens mRNA for KIAA0636 protein complete cds
12669	5.0	AA417030	Hs.5101	Homo sapiens protein regulating cytokinesis 1 (PRC1) mRNA complete cds
1106		HG2981-HT3127		EST - HG2981-HT3127
34367	8.4	AA251758	Hs.40323	Homo saplens spleen mitotic checkooint BUB3 (BUB3) mRNA complete cds
11595	4.8	AA242819	Hs.32539	ESTs
17622	8.4	AA131584	Hs.71435	ESTs Weakly similar to SOF1 PROTEIN ISaccharomyres cerevisiael
34754	4.7	AA287642	Hs.81848	Human mRNA for KIAA0078 gene complete ofs
25038	47	AA010065	Hs.83758	CDC28 protein kinase 2
5312	4.6	U90716	Hs.79187	Human cell surface protein HCAR mRNA complete cds
8085	4.6	AA314779	Hs.105484	ESTs Weakly similar to LITHOSTATHINE 1 BETA PRECURSOR I'M saniens
33656	4.8	W9547.7	Hs.50582	ESTs
8264	4.5	AA401334	Hs.105941	ESTs
34479	4.5	AA262080	Hs.110736	Human bumetanide-sensitive Na-K-Cl cotransporter (NKCC1) mRNA complete cds
7898	4.4	AA263032	Hs.81634	ESTS
10716	4.4	AA053319	Hs.9951	ESTs
2690	4.4	X17620	Hs.118638	NUCLEOSIDE DIPHOSPHATE KINASE A
20203	4.3	N26855	Hs.5858	ESTs Moderately similar to !!!! ALU SUBFAMILY SO WARNING FNTRY III IN sanione1
10923	4.2	AA116036	Hs.9329	ESTS
28050	4,1	AA479139	Hs.75393	Acid phosphatase 1 soluble
10970	1.4	AA129390	Hs.5285	ESTs
4149	. 1.4	U28386	Hs.2397	RAG (recombination activating gene) cohort 1
5767	4.	X53793	Hs.118226	MULTIFUNCTIONAL PROTEIN ADE2
26596	0.4	AA279943	Hs.88671	ESTs
8961	3.9	AFFX-		AFFX.HUMTFRR/M11507_3
38604	3.9	AA598803	Hs.111496	ESTs
30560	3.9	N49284	Hs.1334	MYB PROTO-ONCOGENE PROTEIN
8513	3.8	AA446990	Hs.103135	ESTs
14509	3.8	AA609943	Hs.32793	ESTs .
25284	3.8	AA045074	Hs.110146	ESTs Weakly similar to 52-kD SS-A/Ro autoantioen IH sapiens
27354	3.8	AA425221	Hs.81688	ESTs
18385	3.8	AA227219	Hs.110826	Homo sapiens CAGF9 mRNA partial cds
25240	3.7	AA039713	Hs.110406	ESTs
16854	3.7	AA05552	Hs.71622	ESTs Weakly similar to KIAA0319 [H.sapiens]
3709	37	U07550	Hs.1197	Heat shock 10 kD protein 1 (chaperonin 10)
13606	35	AA456437	Hs.20386	ESTs Weakly similar to CLEAVAGE STIMULATION FACTOR 64 KD SUBUNIT IN sapiens!
8338	3,4	AA417152	Hs.5101	Homo sapiens protein regulating cytokinesis 1 (PRC1) mRNA complete cde

EST - 02869	ESIS Highly Similar to KINESIN-LIKE PROTEIN KIF4 [Mus musculus]	ESTS Basic transcription element hinding protein 2	Zinc finger protein 139 (clone pHZ-37)	NAD-DEPENDENT METHYLENETETRAHYDROFOLATE DEHYDROGENASE	ESTs	Homo sapiens importin beta subunit mRNA complete cds	ESTs	ESTs	Human FX protein mRNA complete cds	EST - RC_AA070364	ESTs	Human translation initiation factor e1F3 p66 subunit mRNA complete cds	ESTs Highly similar to INORGANIC PYROPHC 3PHATASE (Bos taurus)	ESTs Weakly similar to IIII ALU SUBFAMILY		Thymidylate synthase	Homo sapiens MAD3-like protein kinase mRNA complete ods	ESTs Highly similar to phosphorylation regulatory protein HP-10 [H.saplens]	ESTs Highly similar to YSA1 PROTEIN (Saccharomyces cerevisiae)	Homo sapiens RRM RNA binding protein Gry-rbp (GRY-RBP) mRNA complete cds	ESTs	ESTS	ESTs	Human Bcl-2 binding component 3 (bbc3) mRNA partial cds	ESTs	Human (p23) mRNA complete cds	ESTs	V-myc avian myelocytomatosis viral oncogene homolog	Receptor protein-tyrosine kinase EDDR1	ESTs	ESTs	ESTs	CD47 antigen (Rh-related antigen integrin-associated signal transducer)	ESTs	Human mRNA for transcriptional activator hSNF2b complete cds	ESTs Weakly similar to T04A8.11 [C.elegans]	ESTs	Homo sapiens clone 1400 unknown protein mRNA partial cds
	HS.2/43/	. –			Hs.68900	Hs.81690	Hs.8104	Hs.119387	Hs.75801	_	Hs.44131	Hs.55682	Hs.36454	Hs.125123	Hs.25916	Hs.82962	Hs.36708	Hs.3566	Hs.11817	Hs.31730	Hs.73291	Hs.110048	Hs.73596	Hs.87246	Hs.59509	Hs.75839	Hs.3657	Hs.79070	Hs.75562	Hs.107213 (Hs.22595	Hs.15641	Hs.82685 (Hs.62663	Hs.78202	Hs.5080	Hs.139386	Hs.6831
D28589	W3824/ A465342	D14520	U09848	X16396	AA101551	AA181580	H12634	AA463234	US8766	AA070364	W28362	AA094800	F04258	T96690	AA196512	D00596	AA251909	AA489080	AA158132	AA164209	AA278650	AA104023	AA179387	U82987	W93943	124804	F03738	L00058	U48705	AA421164	AA397916	H61476	X69398	AA043944	D26156	AA310967	AA411448	N22895
3.4	4 F	. & 4	3.4	3.4	3.6	3.3	3.3	3.3	3.3	33	33	33	3.2	3.2	3.1	3.1	3.1	3.1	3.1	3.0	3.0	3.0	3.0	30	3.0	3.0	3.0	5.9	2.9	2.9	2.9	29	2.9	2.8	2.8	2.8	2.8	2.8
387	13838	251	3778	2660	17365	33985	19233	13767	4738	17041	15504	7401	18683	23930	11288	170	11659	14134	11140	17925	26530	7445	18055	15174	33620	1932	39556	1605	4536	36200	12313	19867	6081	16708	357	8029	35830	20151

ESTs	ESTs	H.sapiens mRNA for Ich protein	ESIS	ESTs	ESTs Weakly similar to ORF2 consensus sequence encoding and rayance branchism collections.	ESTS Moderately similar to C-I-ETRAHYDROFOLATE SYNTHASE CYTOPI A SAMIT II *********************************	Human retinoblastoma-binding protein (RbAn46) mRNA commissioned	H.sapiens mRNA for TGIF protein	ESIS	Homo sapiens protein-tyrosino kinase EPHB2v (EPHB3) mRNA committee cds	ESTS	Matrix metalloproteinase 12 (macrophace elastase)	ESIS	ESIS	Human antisecretory factor-1 mRNA complete cds	U2 SMALL NUCLEAR RIBONUCLEOPROTEIN A	ESTs Highly similar to UBIQUITIN-CONJUGATING ENZYME E2-17 KD (Drosonnia melanomastical	ESTs Weakly similar to Yel007c-ap (S.cerevisiae)	ESTs	ESTs	Protein phosphatase 2 (formerly 2A) requisitory subunit B (PR 52) atriba isoform	ESTs Moderately similar to PTTG gene product (R. norvegicus)	Minichromosome maintenance deficient (S. cerevisiae) 3	ESTS	Homo sapiens mRNA for KIAA0688 protein complete cds	Eukaryotic translation tritiation factor 2A	EST - RC_R43286	H.saplens mRNA for transmembrane protein mp24	Homo sapiens brain expressed ring finger protein mRNA complete cals	RAN binding protein 1	Homo saptens RRM RNA binding protein Gry-rbp (GRY-RBP) mRNA complete cds	ESTs Weakly similar to No definition line found [C.elegans]	ESTs Highly similar to AAC-RICH MRNA CLONE AAC3 PROTEIN IDictwostellum discondeum)	H.sapiens mRNA for Sm protein F	Topolsomerase (DNA) il alpha (170kD)	ESTs	ESTs	ESTs Highly similar to zinc finger protein [M.musculus]	ESTs Highly similar to HEXOKINASE TYPE I [Homo sapiens]
Hs.62273	Hs.28555	Hs.84974	Hs.5198	Hs.21766	Hs.20183	Hs.44155	Hs.2758	Hs.90077	Hs.97101	Hs.89403	Hs.97101	Hs. 1695	Hs.25282	Hs.27842	Hs.111709	Hs.80506	Hs.5199	Hs.102696	Hs.20922	Hs.5327	Hs.75200	Hs.7487	Hs.82479	Hs.142592	Hs.9877	Hs.81613		Hs.75914	Hs.8932	Hs.24763	Hs.31730	Hs.108527	Hs.19574	Hs.105465	Hs.3378	Hs.48855	Hs.98073	Hs.7165	Hs.20524
AA505133	AA460077	X91788	AA621122	AA490885	AA412528	AA132983	X72841	X89750	AA215333	W73189	AA291259	L23808	AA126719	AA207114	U24704	X13482	H59617	AA187579	AA227261	AA476319	M64929	AA430032	X62153	AA497013	AA487508	J02645	R43286	X92098	AA285277	D38076	AA253031	AA426291	AA600322	X85372	J04088	AA459254	AA357394	T23539	N69263
2.8	2.8	2.8	2.8	2.8	2.7	2.7	2.7	27	2.7	2.7	2.7	2.7	2.8	5.6	5.6	2.8	5.6	5.6	2.6	5.6	5.8	5.6	2.5	2.5	2.5	2.5	25	2.5	2.4	24	2.4	2.4	5.4	5.4	2.4	2.4	2.4	2.4	2.4
28258	8616	6480	14566	14182	35955	17642	6131	6444	1077	42534	34796	1923	10951	11308	4086	5587	19841	7614	11362	13866	2993	12986	5932	38434	38185	1424	21876	6485	1960	452	11701	36390	14420	, 2869	1497	27872	8163	23065	20837

ESTS	EST - RC 051272 s	ESTS	Homo sapiens protein tyrosine phosphatase PIR1 mRNA complete cds	Membrane cofactor protein (CD48 trophoblast-lymphocyte cross-reactive antioen)	Homo saplens voltage dependent anion channel protein mRNA complete cots	H.saplens mRNA for Sm protein G	ESTs Weakly similar to renin [H.sapiens]	Human serine kinase mRNA complete cds	SIGNAL TRANSDUCER AND ACTIVATOR OF TRANSCRIPTION 1-ALPHA/BETA	High-mobility group (nonhistone chromosamal) pratein 2	ESTs	Homo sapiens mRNA for ATP-dependent RNA helicase #46 complete cds	Human 26S proteasome-associated pad1 homolog (POH1) mRNA complete cds	EST - D78129	Homo saplens diphthamide blosynthesis protein-2 (DPH2) mRNA complete cds	ESTs	MYB PROTO-ONCOGENE PROTEIN	ES1s	ESTs Weakly similar to HYPOTHETICAL 27.4 KD PROTEIN IN MER2-CPR? INTERGENIC REGION (Sarcharomyces, persyletae)	ESTS Weakly similar to HYPOTHETICAL 39.6 KD PROTEIN IN MTD1-NUP133 INTERGENIC REGION (Saccharomyces perevisinal	EST - 575266	EST - HG1112-HT1112	Tropomyosin alpha chain (skeletai muscle)	ESTs Weakly similar to PRE-MRNA SPLICING HELICASE BRR2 IS cerevisiael	Isoleucine-IRNA synthetase	ESTs Weakty similar to unknown [S.cerevisiae]	Homo sapiens IPL (IPL) mRNA complete cds	ESTs	Human heterochromatin protein HP1Hs-gamma mRNA complete cds	Cytochrome c oxidase subunit VIb	Coproporphyrinogen oxidase (coproporphyria harderoporphyria)	EST - D28423	EST - D28364	ESTs	ESTs	Human clone 121711 defective marther transposon Hsmar2 mRNA sequence	ESTS
Hs.20990		Hs.4310	Hs.14611	Hs.83532	Hs.7381	Hs.77496	Hs.25863	Hs.75761	Hs.21486	Hs.80684	Hs.71475	Hs.5683	Hs.76887		Hs.103300	Hs.21214	Hs.1334	Hs.27931	Hs.104058	Hs. 10600			Hs.77899	Hs.15313	Hs.78770	Hs.10724	Hs.8130	Hs. 107213	Hs.83550	Hs.83379	Hs.89866			Hs.42582	Hs.5950	Hs.9564	Hs.104558
AA100925	D51272	AA281733	W42845	X59405	AA094989	X85373	AA011134	T29681	M97936	X62534	AA093977	N69352	AA621752	078129	AA252672	AA257971	U22376	N32919	AA251829	AA621340	S75256	HG1112-HT1112	W86469	AA112063	D28473	AA236018	AA113149	AA481403	U28312	AC002115	D16611	D28423	028364	AA609710	W72276	H88535	AA328993
2.3	2.3	2.3	2.3	2.3	2.3	2.3	2.3	2.3	2.3	2.2	2.2	2.2	2.2	2.2	2.2	2.2	2.2	. 22	2.1	2.1	1.2	2.1	2.1	2.1	2.1	2.7	1.7	2.1	2.1	5.0	2.0	2.0	2.0	2.0	5.0	2.0	2:0
17352	28796	26679	24092	5875	7404	6388	25050	41935	3343	5937	7387	20843	28448	651	11688	11803	4046	20276	34370	14582	3461	924	24348	10898	381	11528	25593	38040	4111	61	9112	380	377	28379	24230	40212	8118

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| 12,515 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 11,527 | 1
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	Philipsing building ATPASTP-banding protein (HEAB) mRNA complete eds.	Figure Columnia Compress Cos	MALATE OXIDOREDUCTASE	ESTe	EST: .	EST.	ESTs Westly similar to ORF YORISBW (S.cenevisiae)	£57.	EST - RC_R27975	EST - RC_AAIB0066	Riconscisease L (275-chgoiseademylate synthetase-dependent) inclinior	ESTa	Proprotein convertese subdificients with type 1	ESTs Highly similar to GONADOTROPIN-RELEASING HORMONE RECEPTOR (Bathis movedicus)	Homa sapiens chrate synthase mRNA complete ods	Haspiens mPNA for p0011 protein	EST - U57341	ESTA	EST.	Norman L-kymunerina hydraliasa mRNA completa cds	ESTA	Namen mRNA for apolipoprotein E receptor 2 complete eds		COLIN MODIFICACION SELECTION ALU SUBPANILY : "III [H. repiens]		College Management of the College Management	neran renestre de sprice protect Paris (Paris) mina Complete cus Gent a		EST	HOLL HOLLELTHON	ESTE Weath applicate any contractor as the properties and the properti	EST - RC AATTES	ESTEWARDS IN MOSE 14 PC alement	ESTS	Memo suplens BAC chora RC119C02 from Zo15	Matta metalloprotenass 1 (intersifial collegenase)	Morten mRNA for KIAACO35 gane partial cos	657*	531	thursen Ca2+-dependent activetor protein for secretion mRNA complete ods	EST.	H.depiene mRNA for TRES			CAMP-DEPENDENT PROTEIN KINASE TYPE B-ALPHA REGULATORY CHAIN	n. speed mixta for orbital nuclear bormone receptor	Front appears in a carboning from a factor of the carboning fr	CSTE INVESTIGATION TO NATURAL UNIQUINOUS CALL TO NECOLCT ASE CITAIN 4 (Caenomebolis alegans)	Manual Control	123	PHOSPHATIONINOSCIO	ENT	GI to 5 phere transition 1	Humen brook-IRNA synthetere mRNA complete cos	Obcapon-its peptide-1 receptor	651	EST
3	2000	H2.99231	Hb.14722	Hs 14543	Hs.4770	#.372	Ms 22355	Hs.112759			Hs 12013	HS.119878	Hs.78977	16389	Hs.137991	He.16381		Ha 27835	18.17548	11.01	He 405730	¥ .	75.1232		H 60753		A (5) 18	100659	Hs 2000		Ha. 19736		Ha. 14484	# 22553	Hs 22900	Hs 63169	Hs.75337	H: 62130	Hs 31497	Hs.74574	Hs 36590	19.31748	Ms 24386	27.0		44.576.130	H. 77831		Hs 25570	H: 83319	HE 91447	Hr. 18937	Ha.2707	Hs.109631	Ha. 165	12403	13.97301
70	177514	AA256378	143944	AA055882	AA323787	H20185	247	AA609749	R27975	AA190669	X74967	Roisse	X64810	AA243052	N25086	X81859	US7341	F04022	W67277	U\$7721	AA504482	AA21872	2007	47.00.44	AA054438	KANITZ	MOTEN.	AA521473	AA164289	G3344-HT3521	AAD90642	AA479294	N63512	AA256158	AA132523	X54825	AA065300	AA133250	H93079	D\$7892	HS2702	H42278	11167	44180333	Negative .	A4446486	AA 347183	1,47278	7196CD	W19346	D30037	N39230	X17544	U4D714	501157	238482	AC196060
;	? ?		Š	ă	ě	š	š	2	Š	?	÷	ş	5	ě	ž	ş	ě	5	Ş	ě	9	? !		? ?	ş			2	2	ž	2	•	2	2	?	?	ž	•	÷	š	?	•	? :	:	? ?	? ?	2			ě	ě	Š	ţ	6	÷		D.
68	Ē	28055	4455	10748	=======================================	18383	Z IZ	35570	61517	Tion.	910	41107	288	<u>.</u>	\$ 5 8	6315	4702	19578	24197		2	9	200	707	16835	ğ	23883	30553	11160	1158	2327	37976	8118	11813	10992	5789	16903	7854	70057	24.50	12.01		15 to 1	33880	7007	37004	24835	2168	4362	7334	9127	20814	2635	4368	1533		42.00

7	•			
8 1	2	260555	VC1621.4P	ESIS.
25.75	? ?		11330	Handens mRNA for kinesin- 2
ž			H1 143080	A asolina Nathra M. Dane
787.81			He 12421	EST
10771		AAC) 1346	H 99734	Hame sesions protein chestchatuse with EF-hands-1 (PPEF-1) mRNA complete cds
22.56	91	AA621409	Hs 112988	ESTe
23803	ę	W86423	F. 100.13	£37¢
2222	2	R62831	Hs.28300	EST
3910	ş	AA520709	Hs 20563	ESTs Westly strake to HYPOTHETICAL 90 8 KD PROTEIN TOSHIC 7 IN CHROMOSOME II (C. diegans)
65810	2	AA053636	Hs.129349	E379
29645	9.4	H95840	Hs.42116	ESTe
36405	2	AA428406	H3.10601	Home sepiers mRNA for KIAA0530 protein partial ods
6 229	5.	U21090	Hs.74596	Human DNA polymentse data smed subunit mRNA complete ods
15006	914.	U30248	Ha. 110736	Human burnetaride-aensitive Ne-K-CI cotransporter (NIXCC1) mRNA complete cds
25262	57	AAD44825	Hs.103446	ES1a
1884	ë	AAZ80298	Mt.107313	ESTe
40594	?	N35368	Hs.112227	ESTA
16074	?	AAD01507	Hs 53861	ESTo
27048	ě	AA400670	Hs 79507	Home sepiens mRNA for KIAA0582 profem partiel cds
31484	ę	N63456	Ha 49983	Est's
19304	9	D80832	Hs. 10728	EST.
2000	š	N34636	14. 14.	Hame espiens done 23815 mRNA sequence
3636	ş	AA428353	H3,86394	
39501	914	AA596738	119,109041	_
10201	?	N45124	Hs 6808	ESTA
236	۲.	D13645	12,247	
Ş	5	X14975	Hz. 139305	
15603	9	AA410295	Hs.97911	ESTe
5243	9.	065940	H1 29494	Hunsan brein secretory protein filsecifip (MSECIf) mRNA complete cos
18581	2	140566	Hs.33785	EST
27025	5,	AA609177	He, 109363	
35700	ě	AA410231	Hs,95009	6519
3363	9.4	M54424	Hs. 1355	Cathaprin E
22400	9	R64 109	HE. 29487	ESTs.
70837	ă	N62869	Hs. 107742	EST.
39082	5	AA&20333	Hs.112857	EST
25062	91.	AA017257	Hs.101139	_
40876	ě	N67607	Hs. 109450	_
58.57	š	X58377	Hs 1721	Human mit Na for addpogenesis inhibitory factor
28551	91	H90134	Hs 141382	EST
25529	Ę	F02202	H1 100960	_
6155	?	X74331	H1,74519	
3214	?	U63303	H: 123029	
96892	ě	AA442080	H1.87507	_
400	ş	N49104	Hs.79106	-
37818	9.	AA463954	H 104820	_
23951	2.	AA234556	Nr. 135158	
17077	š	R49406	Hs.28410	
39637	•	HZ6279		-
23190	?	140530	H3.6241	
20131	ě	HE8118	H: 38063	_
24316	ž	AA490500	H1 (8747	
37829	ē	AA470084	Hs 48356	_
22	š	014518	200	-
25:76	Š.	AA621091	78.7	ESTA
41783	2	103637		
13686	Š.	9/96/9	100	TOTAL STREET,
15768	ū	AA100169	48.10348	

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| 1377 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0
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\$?	169100	H-29736	Home sapiens mRNA for TRAFS complete cos
41854	410	876437	Hs.2001	THROMBOXANE.A SYNTHASE
41163	ě	R08176	H5,20118	
8	ž :	AA253400	H: 104326	
23827	9 1	A41144	He. 194768	6314
200	2 2	A4250541	H4.40176	FSTA HIGHER SINGLE IN SOCIETY OF COURT OF CITEMS 139 (9-4)-12 COURTS
86	2	010000	14, 37108	Human MAGE-Sa antiben (MAGESa) cana commiss ests
41752	9	Retoes	Hs,102020	
39302	5	C14944	Hs.12136	ESTA
4689	ž	055786	Te 54485	Human Rev Interacting protein Rip-1 mRNA complete cds
36506	š	AA429810	He.98592	EST.
40204	š	H88296		EST - RC_Heb396
2015	2	E SERVICE	13 30960	Est
200	2 :	AA430725		EST - RC_AA430728
797	2	MAZEZRA	19.0033	Human mRNA for MAA0152 gans complete cds
3 5	2 :	ACTION PROPERTY.	/oci	
1212	? ;	44198719	707.07	
1210		HG17.HG37	2	FST - HC11-1673
22793	2	R96208	Hs 15533	521
36052	š	AA417027	Hs. 104767	_
26574	9	AA279504	Hs 89629	
31818	š	AT18871	K5.2223	Hamo septens mRNA for KIAA0292 gene partial cds
35197	?	AA398120	He:97504	ESTe
17500	7	AA130586	Hs.71331	EST ₃
37979	š	AA479295	Ha 106290	ESTA Highly similar to RING CANAL PROTEIN [Drosophila metanogaster]
7967	9	U6\$437	Hs.05838	Memen homeodomein-containing protein (HANF) mRNA complete ods
28122	9	AA46592B	Ms. 69559	ESTs Weakly similar to LOK (M.muscalus)
26700	į	AA262197	Hs 88002	-
30162	?	AA487501	He,112329	EST.
22593	÷	RYBTT	Ha 29897	
32049	?	AA350857	H 22507	ESTe
1 000	š	H73486	H= 79066	MITOCHONDRIAL 60S RIBOSOMAL PROTEIN 13
2	9	AAG85681	H= 2003	ESTe
37615	ę	AA45B101	Hs.79337	_
7	ŕ	AA127459	Hs. 303788	ESTA
949	ě	H75263	Hs.124151	_
#	š	U45880	Hs 37137	Human IAP Like profesh fi.P mRNA complete cds
26916	÷	AA331293	HS 47378	_
34.68	ę.	AA228030	Hs.120234	
2,5	Ş	AA454610	Hs. 96532	EST
2262	ţ	H46459	700	
00/00	?	M3/827	1120111	_
5.6	è	81.1870	Hs 12246	Human reeds (RELN) mRNA comprete cds
112	?	COMPONINGS		EST - YELDI SOMMSZI
32159	ę	R43153	Hs 95044	ESTI
35.5	Ę	W90709	H 53	Mothe leutemia oral (brie-1) ancagene harnolog
86738	2 :	AA425810		EST - RC_AAA35610
01515	Š	N. SEEDE	49.49268	
	? !	A278878	15.66347	EST.
34014	: :	779744	71700	
762	? ?	MARATO		COLO TENENY BRIDGE DI CICLOCAMPICEPENDENI INANOCAPPICON FACION ALT-0 (N.SEDE SAL
,	? ?	74716-1074148		C21
15474	2	AA417074	10072	
218	?	013540	141,22008	
26100	7	AA242835	H4.5917	Human mRNA for KJAA0281 pena complete cds
41380		R\$4123	Hs 143745	

4777	9	Sign		TOTAL CASES
67.5	?	200		East 100504
	2 9	Water of the		FXTS STATE AND STATE OF THE STA
67.56	: :	Catter		EST
91916	: :			1531
26716		AA262578		
\$1550	9	R33112	Hs.100489	Hamso AF-0 mRNA complete cds
40113	9	H78003	_	531
10201	9	AA0611285	H1 \$972	EST # WOODLY MANDER TO PROBABLE UBIOUTIN CARBOXYL. FERMINAL HYDROLASE HUE: 1.3 [C. 010] mil
37491	9	AA455239		ESTE HONY STREET IS CHROMOSOME CONDENSATION PROTEIN OFFICE (CARROTTED PROPERTY)
23800	:	195729		EST
ž	:	014657	Hs 81692	Human mRNA for KDAA0101 gane complete cos
etas	:	1000		Ubiquên.conjugene enzyme EZH (homokopous to yesst UBC8)
29693	:	H97819	Ht, 42453	EST
26482	:	AA262491		ESTo
23123	=	125306		651
28925	:	AA278392	Ht. 43881	ESTA
0110	:	AAA35840	H 1914	Homo septens mRNA for high mobility group protein MMGZa
X863	•	AA299764	H: 90641	EST
38432	•	051691	Hs 62285	Prospication prigitation from the production of the production of the prospication of the second of the prospication of the second of the production of the second of the
31312	-	H66843	Hs. (36910	ESTS Westery sender to his ALC CLASS of resources for the single sense.
21112	:	R01179	He. 112536	551
31572	:	N71284	Hs. 1 10524	
17903	:	AA160259	Hs. 72354	EST
20747	:	N65842	Hz. 16395	ESTs
£67	:	022200	HB.78619	Homo sapiens tuman permas glutamyi hydrolase (1/2+1) mrCNA control cos
24.85	:	AA251507	Hs.10601	Hamo sepiens mRNA for KQAAD310 protesh perbet cas
38094	:	AA620636	Hs.112264	ESTa
2	:	015126	Hs,36573	Human beta-12-N-acah(ghucosaminy/transferese B (MGAT2) gane complete cds
357.56	=	012164	He.3350	ESTe
7674	:	AA203742	Hs.36957	£51•
4162	:	U31089	HS.121470	Human DP prostandid neceptor (PTGDR) mRNA pardal eda
185	:	047050	H1 24852	Human putative catchan influx channel (http:) mRNA complete cds
W.	: :	AA402227	H15 87345	ESTs Moderately sendes to N-tropomodulin (R. non-epidus)
2007	: :	170802	Hs 23442	Human nuclear fector I-82 (NF182) mRNA complete cds
2.50	:	H50033	Hr 37829	153
1	: :	740145	Hs 21921	6513
47.03	: :	A4000585	He 41175	1,000
200	: :	091004	Hs.101439	1831
20.00	: :	H85434	144 40672	152
1346	: :	M24786	Ha 42993	ESTS MODERNIN SINGE to ITS ALU SUBFAMILY J'WARNING ENTRY ITS (N' sepiens)
7967	:	AA169533	Hs. 72835	EST
21,520	:	R11673	Hs.16957	EST.
Case	: :	AA478917	+ 3482	ESTe Weathy similar to No definition the found (C. clogans)
90539	:	N49072	Hs \$3968	EST
12778	:	W02063	He 55231	EST
26380	=	AA257012	Hs.68054	_
15636	:	X85632	He 65670	Human Ath Inhancior 2 (Ath-2) mRNA complete cds
40812	2	M574.6	Hs.63389	ESTs
200	:	030070	1 8.8	ATL-defred PalA-responsive (APR) peptide
27874		AB7160	Hs.33665	_
40807	:	N62985	Ns.58593	TRANSCRIPTION INITIATION FACTOR HE BETA SUBUNIT
1524	:	M00904	Hs.8037	ESTe
12798	=	R67075	Hs 2074	Zinc fanger protein X-link ad
e publication	: :	AA209467	Hs.97469	_
1960	: :	1447395	Hs 33947	
41607	: :	R57866	Hs 82757	CLEAVAGE SIGNAL-1 PROTEIN
2548	:	M25897	Hs.81564	
	: :	AA232121	-	
!	;			

:::	AA599477	Hs 103345	551. 551.
: :	AA149543.	Ha.5316	
: :	AA020787	Hs.110291	EST. AC. 100.00
=	A18239	Hs 25276	123
:	HD4756	Hs.109645	ESTS HIGHY SIMILED NADH LUBIQUINONE OXIDOREDUCTASE BIT SUBUNIT (Bos teams)
:	O2510-HT2606	_	EST - HQ25/D-HT2606
:	R77778	100	ESTo
: :	4440147		
: :	1118258	H+ 1078	MAC data. It that affects
: =	670322	Hs 82925	Muman transportin (TRN) mRNA connotate cds
:	W14123		EST - M14123 gail
=	AA448357	£.177	6570
:	FIRES	H3.1012Y	ESTe
:	T40891	Hs.6330	£57a
:	WS489S	H1.2164	Connective dissue activation peptide III
:	R07499	Hs. 141384	657s
:	T16282	75.75 EB	WEE1-LIKE PROTEN KINASE
:	RZ6279	Ha 71848	Harten clone 21348 rnRNA sequence
:	H06701	Hs 27948	ESTs Westly sinder to RMOMBOTIN- (PLasplans)
:	R00545	Hs. 18930	EST.
2	O56894	Hs.34782	ESTE
=	AA447666	Hs.77204	Harnan CENP-F kinetochore protein mRNA complete cds
2	AA313387	Ma 133101	ESTs righty sinder to HYPOTHETICAL 64.7 KD PROTEIN ZK1038.1 W CHROWOSOME III (Caenonabalts siegens)
:	AA129547	Hs.81568	#Site
\$	AA02888	Ms.61423	Figure .
:	AA279091	H1.104420	©S1e
;	V07 867	Hs 38842	H. sapiens crRitch for Pitin Lecteta 1
=	W\$3460	H1 91728	Human 75-hD autoantigan (PM-5c1) mRNA complete cha
7	AA172372	He.20008	ESTS
=	195333	Hs. 122730	ESTs Weakly similar to coded for by C. expens cDNA ykt (0p8.3 (C etcgans)
?	X02530	H1 2248	Intentaron (garmana)-induced cell time protein 10 from
~	N70807	Hs 142460	ESTA
:	159905		EST - RC_159505
-	AA252981	Hs. 87599	ESTs Wesky similar to KO7C11 10 gens product (C.stegens)
=	AA996201	Hs. 82702	ESTe
:	AA233795	Hs 85626	
:	13644	Hs 140933	Human mRNA for KIAA0019 cans complete cds
:	735347	Hs 116318	6314
=	AA248864		EST: AA246394
:	A38715	Ht. 123918	Homo septems change 2540 mRNA segments
	AARTHAR	H: 117860	FSTS
:	XTRAIR	He 3155	Control of the Contro
:	. 0.00		
: :		27.07.4	Vienna seriana CTAT Indused CTAT inthibut Ambilia promises of
: :			Company of the Compan
:	3	13.75022	EST E MODELLINEY STERREY OF MUSEUM TRICKS BEINGER STERREY AND
:	H00580	Hs 29889	ESSE.
:	H00615	Hs 24727	E57a
:	AA278925	Hs 88057	ESTa
:	H56911	# 26845	ESTs
:	AA416881	Ha 97383	ESTs
:	AAA49238	118,26838	ESTs
:	AA431486	Hs.107319	ESTs
:	148211	Hs 20854	Homo Sapiens enologensin II recentur nane commisse este
:	WERST	H1 61716	The series of the SAL-1 extremely and the series of the se
:	AA083044	44 (44225	E31*
: :	4411881	607.50	
: :	N39257	Hs 99291	

EST & Highly similar to 605 RIBOSOMAL PROTEIN 1.22 (Rathus nonegicus)	CYTOCHROME P450 VII	ES1.	Cyclin 81	Nuclear factor of keppe fight polypepide gene enhancer in 8-cells 2 (p49/p100)		Romb supers cone zetat manna requesiona FSTs Wester sundar to PROBABLE ES PROTEIN (Human peofomening type 58)	EST	Human (clone 681) Br-cadhain mRMA complete cds	£\$74	ESTS	ESTS WELKING SETTING TO MICALO CLASS O MONORING ESTS IN THIS SECTION OF COMMISSION OF	EST	EST	6571	ESTS DETINANC ACID BECEPTUR RETAU	ES10	Est	Homo sapiens bicaudet-D (BICD) mRNA complete cds	EST.	ESTs	£31	EST	ESTS	ESTS Louis sections alone 71711 sections mONA partial rife		EST: RC N52827	EST						_	_		Colonial considers & sectional and sectional	FSTs Wastiv similar to tehacidine bansponer-like profein (M. musculus)		_	_		E313	_	_	- 7		_		ESTa Moderatory similar to recommend of increasures!		
H\$.54602	#. 1044	H1 14288	Hs 23960	Hs.73090	Hs.91454	2027	E 90.48	H3 44858	Hs.107375	He.109333	He 37380	Ha.40367	H3,96703	H1 85306	Hs.16457	102	H3.98604	Ms.24917	H3.6921	# 742%	H: 99244	Ha 59832	Hr.18917	15.00343	42 58487		Hs 55080	HS.164300	H3.79025	Hs 105289	Hs.57362	Hs.14934	Hs.88739	Hs.75251	Hs. 765	105/60	44.720	H. 65721	Ha 10104	Hs.19322	Hs 48000	Ht 99821	1. 24526	14580	Hs.140859	113 66 153	Ha.12967	Ms.86556	13.73623	147877	Ha 96264
AA129757	X26088	M20598	C20910	608600	N89884	DOCCAN FIRES	AA252537	AA114091	H14969	W85900	C14363	AA456044	AA312551	AA425356	W22015	N51583	AA428633	M87652	M05626	R69218	A4451894	AA002147	H59798	42401	44413134	N52627	WC7583	AA180448	AA199747	AA488814	AA488073	44454747	AA421478	U78524	107963	167710	110022	117063	T56804	AADE8458	N56923	AA620295	131617	806871	844538	F10927	609809	AA415512	AA178845	Reado	AA504307
:	5	::	: :	:	2	: :	: :	:	•	:	: :	:	:	:	: :	: :	: :	I	፡	÷	:	:	5	: :	: :	; ;	: 2	: :	: :	:	:	2	62	2	7	= :	: :	· =	-	:	:	: :	: :	: :	3	3	:	:	2	-	: 2
23675	22	2000	3933	977C	1631	2000	ž.	25599	19749	4258	146.17	27131	288	27380	200	2002	Z X	8738	39070	22097	37308	18101	20829	8 3	2	10000	22882	16072	16231	38282	28125	17464	366 te	2062	ž	42103	1000	8	23335	10067	30663	14528	i i	21748	21840	9062	18774	22/20	18062	27863	1973

£51.	15 T	ESS MATTER MRNA for KLAAD185 come cardial colo	Numera anticudant enzyme ADE37-2 mRNA complete cds	EST - HG3548-KT3744	EST	16.73 - 16.73 - 16.73 - 16.73 - 16.73 - 16.73 - 16.73 - 16.73 - 16.73 - 16.73 - 16.73 - 16.73 - 16.73 - 16.73	ESI.	EST - X98256_cds2	EST.	E37s	E314	Humen mRNA for KUAAD197 gene partial cds			Human platefet (actor 4 varabon 1 (PF4var1) pers complete cos	EST a Western Growth to pudative price [M.sapens]	EST - RC AA487495	Numen chromosome segmention pene homeloo CAS mRNA complete cds	ESTo	Rethal plannent epitieskum-specific protein (65kD)	ESTs	SRY (res-determining region Y)-box 9 (cemporastic dysplasia autosomal sex-reverset)	EST.	Hamen mRNA for KIAAD029 gene panial cds	1950 Martin services martin. J press can to Delicity and a services and	MOTO september those 27 m SNA shannesties make make waters when a necessary of	ESTs	. EST - X82279	531	Hisablens mRNA for Michaele phosphoprotein myc9		E314		CDC21HDMCLOG	ESTA	Homo septens done 23622 mRNA sequence	Human hundingth Interacting protein (HIP2) mRNA complete ccs	Hydroxymesty/Dane symblese	and the control of th	THE STATE OF SPRINGERS OF SECURE OF SECURE SECURES CONTRACTOR CONT	Peruzinamel membrane protein 1 (70%) 2 elements	ESTA Moderately senden to M-chare phasebooker 11 ft seniens	ESTA	£51s	ESTs	£51	ESTS Workly strains to LINE-1 REVERSE TRANSCRIPTASE HOMOLOG [H. sapens]	531	101	EST: WXX01	E31e	E31s
		Hs 45024 H			75.4230 E.		45.109270 E	ŭ	11.144000 E	HEMSAT E		Hs.22559 H	_			4 106937 F.		H: \$0073 H	Hs 55378 E				_		H: 24395 E.	_	_	_		-	_	H159477 F:	_	_	H1.105576 ES	Ξ	_	Market M		_	_		_		_	-	_	18.27.34 ES		ט מ	115.84497 ES	
	AA262179 H			7	110248		_	X36,766	Z38612 H		_			_	M2016/	Ī	_						_	CLESTS				X82279				4 (2)(8)		X74794	_	_		7.70034					_		_			H0/084		H30201		AA428137 H
:	::	: :	:	:	: :	: :	:	:	:	:	•	=	3 :	= :	: :	: :	-	?	•	;	ç	: :	: :	::	: :	: =	:	3	:	*	:	: :	:	:	;	:	•	::	: :	: :	:	3	Z	Z	;	I :	2 :	2 :		: 2	: :	:
20550	28470	8	683	25	2238	¥1.2	25.822	6602	42701	Z 252	18290	ĉ	8 5	000	2 5	\$2000	33180	7727	22920	1785	24673	9289	2000	207.	22528	17/04	31596	6259	31578	13207	ž	11592	26843	6160	37887	42518	Ę,	,	749	2500	7112	11257	31467	26924	34928	50662	10925	18081	M70	Ą	29822	27411

6516	EST.		EST - HG2848-HT2863	£37s	ESTs Weekly similar to unknown (S cerwises)	Human small profine nch protein (sprd) mRNA clone 1282	63%	ESTA Weekly survivar to SPERMATIO SPECIFIC PROTEIN 12 [Secre officinels]	EST. AAJOSIIB	ESI's Weakly strater to week sindantly to procokagen agota chein 1(V) chan (C. etagens)		ESTS WARKLY SETTON TO HOSACOAS JAN. THIS GROUP	ess.	Haspions mRNA for apoplasis specific profess	Pregnancy-specific bate 1 photoprotein 8	Est ·	est :	EST	Problemang cas nuclear anagen							EST	EST	Hespiens mRNA for M-phase phosphoprofen mpg9	EST.	ESTA	ESTs Weakly similar to MOESIN/EZRIN/RADIXIN HOMOLOG TO malanogasted	Eukaryode transladon indaton lactor 3 (ell-3) para suburti	Homo Espans metha na Puladad projet comprete cas	E.O.1 Money envises revises manufulms published; a (PRCI) mRNA complete cds			EST. AAA81218		_	E01s				_	_	_			FESTS CONT. MALCOL. Charles to companie accordance contain second 1 10 melantomation				- ESI*
Hs 47076	H3 60556	787.4		H1 495 10	Hs 25740	Ms,2421	Ha.83842	£ 20		Hr. 126781	F. 104	Ms.38399	4, 97.16.	121	Hs. 108938	Hs.47338	Ha.98023	H3 44610	H3.70996	Hs.44690	100263	20000		H. 17830	He 98003	Hs 07529	Ha,96330	Hs 86178	Hs.12544	Hs 42979	Hs.35176	2000		4.610	H3 47476	Hs 22970		H1 27621	£ 36573	# 49380	H. 6065	Hs 20137	1	Ha.106794	Hs 79440	Hs. 101619	# 73.29	110 99210	C0C21 9H	11.673	144123	H. 73606	Hs 94161
N50556	AA599309	C20532	G2646-HT2983	F06925	AA186304	M21539	AA287450	H97562	AA305116	N20583	AA210722	AA250171	2001	C21245	X17098	N51761	AA406167	AA256460	M15738	N35065	103886	*******	7,000	F04134	AA408083	AA399053	AA435750	AA43686	AAA43528	AAB18145	AA405098	19678	WEEEE	***************************************	. 079797	N\$2189	AA481218	AA292436	015128	AAD04420	4476781	AA787567	140589	AC195N	W67008	840CDH	AA129395	AA449121	F04877	AA777803	822183	H11360	NOOGS
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\$1905	26313	16321	100	19578	11232	2486	26643	40331	g	28783	8	8012	19761	28389	56.84	01700	35765	26360	<u> </u>	30262	41702	200	2000	77.01	15746	35356	36769	36900	27555	16290	27113	700	200	6007	70701	800	8778	M028	14965	18115	96.5	14870	23211	1004	11927	338652	17581	3723	16712	2005 2005 2005 2005	27.5	167.00	31295

24K7	;	239106	Hs 27285	100
31292	;	N66615	H8.49232	ESTs
1285	;	G4157-HT4427		EST - HG4157-47 432
138	:	G2981.HT3127	~	EST - HG2981-HT3127
16212	;	AA198506	Hs.66076	EST
78.7	:	AA251756	H3.40323	Name saptens spleen missuic checkpoint BUB3 (BUB3) mRNA comolete cos
34602	:	AA291458	Ks.98504	ESI
34762	=	AA287834	Hs. 102909	ESTs
1395	;	AA242819	Hs 32539	551,
232	;	AA405082	H3.125014	ES1s
17622	;	AA131584	Hs 71435	ESTs Weekly similar to GOF1 PROTEIN (Seccharamyces cerevisiae)
12.5	÷ '	AA408335	14 95 187	E37s
7 7	=	AA287642	H3.61248	Human mRNA for KAADO78 gene complete cds
2223	•	147291	Hs.8610	EST
37667	•	AA460318	H: 10165	EST's Highly similar to 60S RIBOSOMAL PROTEIN LS6 [Radus non-egicus]
136	7	AA236786	1 20076	EST:
39622	:	AAS98967	Ms. 141982	EST
7118	=	079296	13 74842	Dihydrdipoemide S-erchitansierase (£2 component of pyruvase dehydrogenase complex)
25032	÷	AA910065	H3 63758	CDC28 protein kinasa 2
19289	Ç	H16567	Hs 21253	EST.
10520	÷	117045	Mt 90283	Collegen type I alpha-2
121	-	K#055	2,5	SOOUM CHANNEL PROTEIN BRAIN & ALPHA SUBUNIT
8	Ţ	13861	H4 69650	kniegraf Vensmembrane protein 1
35400	÷	AA389591	Ha.97831	Homo suplens putetve DNA methymensierse (DNMT2) mRNA complete cas
35248	7	AA398367	Ks 97617	EST Weakly similar to HSP80 protein [M.musodus]
200	Ç	AA426270	1, 36498	ESTE
21509	Ç	R27314	Hs 23820	
31381		N67838	Ha,49397	ESTs
22/22	÷	AA2827781	H1 95321	ESTs Highly similar to PRE-MRNA SPLICING FACTOR RNA HELICASE PRP28 (Seccharomyces cerevisiae)
36336	:	AA425151	He.79310	Human GAP SH3 binding protein mRMA complete cds
17409	÷	A113138		EST - RC_AA113138
8.	:	U67156	Hs.B4149	Muman natogen-activated thase kinase kinase S (MAPKKS) mRNA complete cds
100	;	N49967	H1.48624	ESTa
252	;	AA489847	Hs.1(2019	ESTa Weathy similar to LINE-1 REVERSE TRANSCRIPTASE HOMOLOG phycicabus coucangi
200	;	44433950	Hs 22530	
6	;	N21814	#8.1003S	Homo septens besto-leuchte zipper trenscription fector Mario (MAFG) mRNA complete cat
1477	;	AAS09427	45.24164	ESTS MOGRAPHY BITGET TO BIT ALLU SUBFAMILY SC WARNING ENTRY IT! [H arapions]
36213	;	AA488347	75.11380	ESTs Weakly similar to putative p150 (H. cepiens)
212	;	90716	13.79187	Numan cell surface protein HCAR mRNA complete cds
\$228	:	W70326	1550	E57:
97.54	: :	2000		
200	: :	AANSTORE		CCT AAARTON
2512	: :	141819	H. Poun.	Home series roots beneate blace founds. (FDMP) moust complete out
37066	;	811519	Hs 52054	EST
2008	;	AA314778	X 105484	ESTA Weakly stratur to LITHOSTATHINE 1 BETA PRECURSOR IN servinal
727	;	013633	Hs 77835	Human mP.NA for KIAA0003 gans complets cds
34006	‡	AA186761	Hs 80961	DNA роўтаства датта
33656	ŧ	W85477	Hs 50582	EST
24045	:	AA105517	H3.11150	ESTs Weekly similar to 111 ALU SUBFAMILY J WARNING ENTRY 111 [H. sepiens]
6028	=	X86503	Hs.90011	Adenytosuccinete synthese
8 1	•	029483	H3 108162	Cytochrome B361
70704	: :	700000	HS. YORK	201
18922	= :	R85208	13.74.16	EST
ĝ,	;	M33969	HS 91107	A saperts thank for the storogen
- C	: :		1001	
2000	: :	44413047	12 57 57	COIL PRE-
2442	; \$	AA262080	H4 110735	Colta Numerical sections Na-RCC commander (NKCC1) mBNA possible and
15921	:	Y12065	Hs 5092	Here segions michael content and the northeast physical physical property of the content of the
•	•			

11279	:	AA195399	H3.24641	EST
27.53	: :	AA621348	10042	ESTS Highly smiles to DOLICHYL. PHOSPHATE BETA-GLUCOSYLIRANSFERASE (Saccheromyces ceremines)
17.78	: :	AA481168	H- 67777	2512
22153	: :	851401	Me forent	Months and the management of the second of t
7894	;	AA263032	H 61634	name argents menne com dispressione 5421-22 done ASA
19902	:	#56738	£ 22 t8	E513
9276	;	D62374	Hs.131854	E37s
10716	።	AA053318	Ha, 9951	EST
13193	;	AA442783	H3,20483	ESTs Highly similar to 02/MITOTIC: SPECIFIC CYCLIN B2 [Mesocycelus suralus]
8	:	X17520	Hs. 118638	NUCLEOSIDE DIPHOSPHATE KINASE A
15102	;	44371509		EST - RC_AA171509
17863	;	AA169226	Hs.72762	EST.
24962	\$	HUMTFRRAM11507	1907	AFFX-HUMTFRRAN1507_S
31680	:	17413	Hs.50452	ESTs .
27.168	:	AA410258	Hs. 85908	ESI
17.07	•	020381	H2 92453	
26348	;	AAG08752	£ 1989	EST
er e	: :	AA018387	Hs 40515	ESTS Weakly sender to MIALU SUBFAMILY SP WARNING ENTRY III [H. sepvens]
900	: :	W48500	Hs 39972	ESTe Weakly similar to transposon LRE2 reverse transcriptions homolog (H. appara)
1000	: :	A35164	1	
96.56	: :	40000	11.780	COLO WORTH ATRICTO ET ALU SUBFAMILY SO VIARNINO ENTRY EN PLASPONS
20,52	; ;	AAARSONA	H. 1704	ברי. הריי
38690	;	AA600121	Ks.111467	
20203	;	M26855	Hs 5656	ESTS Moderney similar to 111 ALU SUBFAMILY SO WARNING ENTRY 1517H segions!
10251	;	H 78:185	H3.18171	ESTs Weakly similar to CO IHG.7 (C. elegens)
12684	÷	AA417558	H3,25208	EST.
31636	;	N73680	Hs.57435	Natural resistance-associated marrophage protein 2
20789	;	N67277	8	
7761	: :	No local	1	EST - KOTESA
K	: :	AA252414	104301	EST.
10132	;	R35733		EST - R15733
16629	7	AA036811	Hs 61859	EST ₃
25148	;	AA026356	Ms. 105108	EST
28730	:	023959	Hs 5858	ESTS Moderalely similar to 111 ALU SUBFAMAY SO WARMING ENTRY IT! [H.sepiens]
10200	7	125794	H9.77361	EST.
2699	?	AA600178	#8.1274S	E31s
	:	055794	18.489Q	15319
900	: :	666/CM	He 100018	EST6
1	: :			econoci i sonoce
i či	: :	AF006442	1 400	numen reconstruction advants protein (RAG-1) gene complete cos
13.79	;	AA449741	H3 4029	EST Weath circles to A. & PROTES IN account
4.2	;	U78293	Hs 90002	Munan done 23948 mRNA sequence
2832	7	19585M	H.2077	Zint Brigar protein 8 (close HF.18)
38005	;	AA478959	Hs. 105624	E31s
36573	።	AA431095	Ma.96706	EST
18290	:	AA213620	Hs.46301	ESTs Weakly sinder to putetive p150 (H sepiens)
Ę.	: :	H63953	:	EST - RC_H86953
2	; ;	ALCINION A		Arratum resount 130/_5
25.00	: :	20113144		
573	Ş	H52038	44.26710	TEST
11/07	Ş	N\$3564	H1 100159	
\$;	U28386	Hs.2387	RAG (necontainstion activisting gene) cohort t
5767	:	X53793	Hs.118228	MULTIFUNCTIONAL PROTEIN ADE2
e e	:	X05232	Hs,83326	Stromelysin
20210	:	N34893	H-6153	ESTs Maky similar to HYPOTHETICAL 47 B KD PROTEIN B0280 9 IN CHROMOSOME III (Caenomaddiss elegens)

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| 1971 | 1970 | 1970 | 1970 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 | 1971 |
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| 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100
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EST - AA142144	_		_	_				-	_	ESTs	_	_	_	- 101		-	-	EST	ESTs	•	_	_				FOUND ASSESSMENT LINES OF THE PROPERTY OF THE					ESTS Westly similar to GA BINDING PROTEIN BETA-2 CHAIN IN sapiens			_					-		_	_	Homo Sepiers (done 81835 13) chromosome 4p16.3 DNA fragment			. 63Ta			_				Harman I DN centern miDNA controlle acts	
	Hs. 109156	45,87157	Hs.5122	Hs.104135	13.7656	1000	H12170	Hs 6309	Hs.53810	Hs 21738	H1 24287	Hs. 125123	Hs 46495	100	7.00	Hs 20787	H1.0179	Ha.54174	Hs 25916	Hs £2962	Hs 63272	H# 49915	3670	77.7	10211.01	3	Hs 25245	100043	H3 53149	Ha 103045	Hs 63220	He 101571	Ht 27149	Ms. 131361	110 48269	K. 3709	H3.62394	15140	10033	M. 5337	He 11817	Hs 32753	Hs.94799	Ht 27693	Hs 43100	Hs 30619	Hr 66870	H1.23383	Hs. 58197	1.082	H3.48842	R/IS S		Hs 71814
AA147144	R78248	M92438	U41387	AA206983	WEBERS	44791921	M80627	AAA53444	H26011	R15846	AA280928	19690	N45276	Cleary	44427579	R60367	M66818	N64408	44195512	000396	85050	AA243133	AA251909	HIOPER	06762944	440047	AAABBBBS	R55078	AA102566	W42928	194828	R78618	M35999	M&6606	AA112879	058253	128021	***************************************	W45c91	W78050	AA158132	C20679	134409	AA610108	NZ2162	221420	X95632	AA443811	AA424038	AA456821	AA412429	A 164 209	1,4000	A14724
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26530	2 2	AA276650	Hs 24639	EST
9099	•	AA053258	Hs 3/821	Home septens mPNA for KIAAD648 protein pertial cos
22.00	: :	110277	Hs 115873	Monte suppens ribonacteese P profess subunnt pall (KPP40) gene compete ons ESTs
33585	: :	W93000	Ha.59389	6373
ន	•	013627	119 64021	Human mRNA for KIAA0002 gens complete cds
4298	2 :	036448	Hs 74574	Human Ca2+-dependent activator protein for secretion mRNA complete cds
(Page)	2 5	AA104023	H9.110048	
(8055	: :	AA178337	Hs 73596	
7262	:	AAD83339	HS.126781	531
8748	:	H03666	Hs.112013	ESTA
ğ	9	087009	Hs 43624	Human (lambda) DNA for immunoglobin fight chein
38447	ô	AAS04255	14 SA 204	Human protein tunese ATR mRNA complete cds
4 64	2	R46837	H2 107450	
8	2	Libiai	12134	TRANSLATIONAL IMITATION FACTOR 2 GAMMA SUBUNIT
2776	2 :	4444039	1.00000	1381
5 5	: :	TATOM	9,76	The Sport contain & APEN
15174	: :	U\$2587	H1 87246	Human 8642 binding comeconam 3 (bbc3) mRNA parties cds
23620	: :	W93943	Hs 59509	6571
1419	•	Y11661	H19.9904	Home sepiens ribosomal protein 512 gene nuclear gene encoding misochondrist protein complete ods
41077	2	N95028	Hs.125031	E31,
1932	2	2682	Hs.75839	Human (p23) mRNA complete cds
39556	ç	F03738	Ha.3657	£31s
15108	•	AA002258	Hs 59939	EST4
32156	:	R40381	HS, 147852	637.
1361	:	AA456648	Hs 20061	
-1949	ç	AA281251	13.000	ESTs Weakly shriker to inflormy protein that (D metanografier)
800	2	X68194	H3.60919	Pentophysis Duman temitrocyte fine NeCe1 mRNA 2108 ng
2	2 :	W27374	H 2300	Hartes septens (OkD protein (BC10) mR(NA complete cas
38080	٤ :	AA482337	10013	E31
200	. :	A44/0004	1	
50.50	: :	or serve	48007	H. separa P.A.P. mRNA
4 5	: :	028/21	E	Prosperior to grain data A (peroxy arm) nocularia inemograciana)
1	: :	Water.		
1007	: :	44331677	M. 04777	
200	: :	101101	2000	
29249	: :	HS2918	H1 39165	E513
947	:	U99815	Hs 85813	Human SMIGNF complex 155 KDs suburit (BAF 155) mRNA complets cas
10104	=	R23855	Hs, 107968	E31.
15039	:	046116	H1 89627	Protein tyrosins phosphalese receptor type gamme polypeptide
ş	:	1,00058	He 79070	V-myc evian myelocytomatosis vital oncogene homolog
\$ 29	•	048705	111 75562	Receptor protein-tyrusine kinese EDDR1
£550	:	R38676	Hs.102963	ESTs Weekly similar to cell division control protein COC21 P1 sepans
26555	:	AA279071	11.94.945	ESTs Weetby similar to TD8A11.2 [C.steguns]
104	€	U41515	H9.65215	Human delated in spirit hand/spirit font 1 (DSS1) mRNA complete cos
21009	7.0	NSO403	Hs.28928	ESTs
3602	:	716105	Hs.117848	HEMOGLOBIN EPSKON CHAIN
4833	:	U63455	H3 54470	Suffornjúvnes mosphor (hyperfinauknemis)
36200	:	AA421154	113, 107213	
362	=	AA281078	Hs. 109221	E516
38299	=	AA398622	Hs 73133	
og :	= :	M74558	H3, 142363	Martin Sit micks compared cos
2010	::	44747018	4. 22.445	PATE
5928	: :	XETCAR	Ha 75185	WEELLIKE PROTEIN MASE
SZEC Movel	: :	600155	1,2827	

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A13798 H 4115 EST MOSERNY JEMET IS CHITETRAYTOROFOLATE SYNTHASE CYTOPLASMIC (M. 1849ens)	*		0 Ht 142190	Hs 4214	_	AAAAA UAAAA CAAA	H: 33977	Ha 90077	XB1072 Nr.92996 Hurren mRNA for T and receipts close 10RA:17	X78627 Hs.75056 H.sepiene mRNA for bunsin		Hs 97101	Ws 22587	WASTER HERSON PHOTO EASTERN SPOREST TO THE SEASTS EPHDEY (EPHOS) MRNA COTROLES COS	H1 95265	Hs 107323	44,73103	148,77493	AAABASON NASTASI ESSTE BARRING LALACEME ETT	Ns 21603	4 Hs 7523	AA441720 Hb.7551 ESTa	H3,3430	Hs.101368	1, 1003		M. 1933 M. 16.03 Millerin Roching Day C. S. Company Company Company Fitted Fitted Millerin Roching Company Carlo C	9 Ha 97101	Hs 3281	Hs 1904	H1.0768	CR84-HT684 EST - RC864-HT684		13604	Hs 56013	Hs.107914	3 Ha 77719	DE2775 His 10724 EST's Wealty strate to unknown [S. carefulled]	•	H1 1814	4 Hs.109622	L23805 H3.1695 Matrix metaltoproteinese 12 (macrophege éssisse)	Ha,6817	Hs 46572	H 97721		ACMING HEALTH FETT.	H9.37573	8 Hs 23170	Hs 29405	WS5015 N4 34620 ES14
: ;	: 2	~	23	5	: ;	: :	: :	:	7	~	7	2.	~ :	::	: :	:	7	2	2 2		2	7.	?	2	2 :	: :	: :	~	2	2	2	: :	: :	: 7	2	11	23	⊋ :	: :	=	:	"	2	:	: :	2 ;	: :	:	*	. :	2
ž <u>š</u>	41429	17052	34263	22437	2002	16241	100	Ī	5316	6240	42116	Ē	200	100	36256	8181	14769	32961	35273	2583	25.25	13723	Z		8 9	3 2	90691	87.7	41955	200	33686	COE.	76.70	107	2125	40528	25245	2674	7635	35	7363	1923	24906	2000	P. 1	2007	8 25	19022	1727	22010	216

22

9554 18104 24582 40038	***	H81487 AA168801 241563 H69485	Hs 12063 Hs 85634 Hs 26975 Hs 6236	÷ 5 5 5 5
22148	: :	A9002359 R51831	Hs 105478 Hs 25829	Human mRNA for KIAAQS6 I gana KIAAOX61 protein EST19
<u> </u>	: :	US1990 AA426156	Hs. 94178 Hs. 42714	Muman hPm 18 mRNA complete cds
22402	3.6	N49300	Hs 24906	
41807	₹ :	R67258	Hs.58215	ESTs Moderales similar to nucleatin (M musculus)
, G		1962501	# 78.83	Munan matan in MAGNATI of gene partal con Munan penjasas Manda Carradala cola
16907	. 2	AA053298	Hs 53136	ESTs
15268	:	W07562	Hs.26196	ESTs Moderately similar to rAS (A nonegous)
38623	2;	101716	Hs. 105153	ESTS.
10951		AA(26719	Hs 25282	KOLF NIGHT BITTER TO DITTER THE FAME THE FAULTH IN CHASE-THATE THE EXISENT REGION (SECTIONARY CONTINUED). ESTS
6150	2	X74262	H# 81058	RETWORLASTOMA BINDING PROTEIN PAB
26136	:	C20945	Hs 108117	ESTS Highly similar to HYPOTHETICAL 34.2 KD PROTEIN IN COC12-ORCS INTERGENIC REGION (SEKOLAROMYCES CARANISES)
28831		AA29282	12 13 13 14 14 14 14 14 14 14 14 14 14 14 14 14	eSile MgNy santist to moduletor recognistic lector 2 (N. septens) ESIs.
2173		142621	Hs 83062	Home seeins 1 v. 9 mR IA control in cos
10642	: :	44040149	H#, 110103	Human Circumosome 18 BAC clone CIT9825K-A-270G1
15026	*	041815	Hs 91161	Human C-1 mRNA complete cds
7699	:	AA215299	¥4.70830	Horno sepiene chramosanne 18 coemid R36781
Ž,	:	X858X	Hs. 112743	Home serviens mRNA for SCP-1 complete and
300	: :	N62122	1000	2013
900		2470	Hs 21/708	E.C.1.5 Human antite econtrary factor: 1 mRNA coversis to cits
20015	2	AA598938		EST - RC_AASBesis
11619	:	AA258189	Ha.32471	6514
37433	:	AA454103	He. 1 10031	ESTe
28270	2	AA521188	F. P. 217	ESTA
3567	2	X13482	F1.6050	UZ SWALL NUCLEAR RIBOANCI EOPROTEIN A
200	: :	HSMIT	Hs.5188	ESTE HIGHY INTERED URIQUITINACONTUGATING ENZYME EZ: 17 KD (Drauchka melanogaster)
14053	: :	AA485147	Hs (2763	ESTS HORN similar to HYPOTHETICAL 516 KD PROTEIN ROSDIO 2 IN CHROMOSOME IN Classocherblica alegests
31874	: :	M171M	100	Triban tanoanous i i moronous a vincous con toto toto toto toto toto toto toto
7614	: :	AA197579	102598	ESTs Weathy stratur to YaXOO7c.ep (\$ certainties)
17878		AA479195	Ha 105620	ESI
7090	~	44009913	H1 103300	Momo sepiess depthemate bissynthesis protein-2 (DPH2) mRNA complete cds
17852	2	AA196360	£ 7623	(51)
19070	: :	H05970	Hs 133028	ESTE PARTIES CITIES TO STAND T
1771		44136569	H1 144295	ESI
39969	2	AA599694	Hs \$7730	Hyman mRtA for KAAD111 gene complete cds
70867	:	H79565	13.2962	£57¢
8 3	: :	031448	H2 10438	Horno sapana breskoari duster region protein 1 (BCRG1) mRNA Comprete cos
1 2	: :	AA459555	H4 3/873	Horno series mBNA for KIAAIMAR worden contist erts
13956	:	AA476319	Hr 5327	EST.
10303	7.	R96178	Hs.51167	Atacia laisangiects sia mutased (includes transfernantsion groups A C and D)
64272	2	R59601	Hs 25679	
1829	: :	44206391	10,100	EST: WC_AXZ05391
38552		603603	He 75574	ESTATATO RAS RIBOSOMAL PROTEIN
27530	: :	AA435999	Hr. 134132	6516
1785		13434	Hs 64162	Harten chromosome 3p21.1 gene sequence complete cds
14746	3.	76007	H\$ 60315	HAMMIN MRINA for KLAADOOT gene perfei cds
£	3.0	M64929	Hs.75200	Protein principhatase 2 (formerly 2A) regulatory audumid B (PR 52) aithia ractorm

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18003	•	W171592	H1 70980	£51s
24198		W67524	H1 82811	Murtan protein-tyronine procephalase (HIU-PP-1) mRNA paries sequence
78.		W92703	H3 100239	EST
949	•	96/952	Hs. 142200	EST Weekly amiliar to putative pt 50 (H. septiems)
200.00		7 X	113 40680	ESTs Weakly shriter to CSOF4 12 (C.stegans)
666	_	W(2)483	Ns. 26676	ESTS Weakly similar to probable CBP3 proving homolog (C. elegans)
50103	.	20/50/00	H3 /68//	
			2011	ESIa Moderately activate to ZMF127.XP (H suprens)
		M2/300	20/20	ES1:
	•	200000		Warnan kinesin-tike spiridle protein HVSP (HVSP) mRNA complete cus
5000		MONEY OF	100	ESTS Weakly similar to LINE-1 REVERSE TRANSCR:PTASE HOMOLOG IMycticathus coucangi
	•		13.11045/	
2/7		200	F6. 1027.70	651
96.01		0292	77030	6510
		M3ZZCH	47433	ESI
77		20.00	78328	Home septens protein phosphelese 2A 836-spison (PP2A) mRNA complete cots
6091		W24303	Ms 21167	ESTA
970		1633	18 78008	Decargoymidyfate kinase
01221	•	7283774	Hs.21265	ESTE WERLY SIMAL ID PROBABLE TRYP! OPHIANYI, TRNA SYNTHETASE MITOCHONDRIAL (C DEGRAS)
700		2000		EST - S63004
/047		M44758	2 2	Homo sapiers neptvocysin (NPHP1) mRNA partel cds
7,070	ς ;	10774	18.20530	Harman mistrak tor kinesin retisted protests partiel cds
7			1	Covin Bi
2/00/2	•	2000	200 F	551
i sara	52	000	H 2690	
17269	~	A085179	Hs 24382	EST
8000	:	22.5	H3.11449	ESTS Highly similar to MYPOTHETICAL 30 1 KD PROTEIN IN APEIALAP4-CWP) INTERGENIC REGION (Saccharamyea caravasae)
Z S	•	X62153	H: 62479	Mitchnomosoms maintenance dehident (S. carensias) 1 -
15685	2.8	X85073	Hs 96747	Hisapiens mRNA for vendin essociated protein X
17852	5.2	AA165677	Hs 65757	ESTs Weathy similar to F18A11.1 [C.elegans]
12187	·	AA283208	Ha. 10852	ESTa
6210	\$2	X76942	Hs.77335	Marto septera polgin-245 mRhu complete cds
34047	٧ ٢.	A194156	Hs. 106928	ESTs Weathy similer to G1 10 S PHASE TRANSITION PROTEIN 1 HOMOLOGIH saciens
16929	*	AA058852	H* 62590	ESTe
26934	, s	W287138	Hs 59746	ESTs Weathy similar to ASPARTYL. TRNA SYNTHETASE Thermus equations the minimorphical
5157	5 2	120031	H1 66563	Human mitochendrial intermediate percensor (MIPEP) mRNA mitochandal cene according mitochendrial puries and
XXX	¥ 52	£1078MA	HB.142597	6570
13761	•	W72987	H1 58257	ESTs
26991		AA398284	Hs.48050	E5Ta
1380		AA17350S	H. 35353	ESTO Woody simbly to hypothetical 46.1 kg protein in PhO2-POL3 intercenc region (s.comyniao)
14.00		162507	H* 89372	Mumes fetal Alt-50-reactive ctone 1 (FAC) mRNA complete cds ·
1363		AA455899	H1 22151	ESTS Highly, similar to HEUROLYSIN PRECURSOR (Sus across)
1086	•	A412151	Hs.108974	
20.00		AA487503	H2.9877	Harra expiens mRNA for KUAA0581 protein complete ads
100	•	W284744	M3.75510	Annexin XI (36kD autoentigen)
1424		202648	Ha. 81613	Euraryokic translation initiation lactor 2A
16778		A404700s	Ha.62900	EST
21576		R43286		RC_R43266
	•	×1004	£ 38.88	EST
22.20		238588	H. 7903	ESTe
1		M242904	Hs 40837	Hamo septens profits rich Gie prosein 1 (PRGP1) mRNA compuste cds
7474	•	A126582	8	EST a Westly similar to No definition line found (C elegans)
24780	_	W236868	fb.111314	ESID
5316	-	206060	Hs 79365	Human done 23374 mRNA sequence
10218		R68654	Z .	ESTS MIDNY SIMILES MYPOTHETICAL 44.2 KD PROTEIN IN SCOZ MRFI INTERGENIC REGION (Saccharmycss Grenssas)
81.00	; ;	X155981	11.50169	Home supports retrocks shown as accepted protein HEC mRNA compass can
ŝ	2	x85086	41 75914	H septens mRNA for transmentarine protein mp24

23.55	2 2 2 2	AA342959 W74751 AA430160 B10301	Hs.110041 Hs.42765 Hs.20584	EST : EST : EST WINN'S INDIA IO FZSHI I COMPANI
0250	2 2	A:006265	H3.8222	E.3.1 Homb sapiens cancer associated surface analogan (RCAS1) mRNA complete cds
30037	*	N27439	He.51652	ES1s
27602	?	A4443702	Hs.29635	ESTs Weakly similar to W02812.7 [C.elegans]
200	Ξ:	559184	Ha 78350	RYK receptor-uka hyrosma kunasa
200	: :	**********	H4. 14850	501E
12.00	: :	H24000	H3.93048	Memo seoiens DAC clone RG300E22 from 3e21-q31.1
29392	:	H72914	Hs.103318	ESTe
35521	:	AAA00831	H1.111916	ESTe
20324	:	N35406	Hs,74014	Phosphologis C bets 4
16620	٠.	F02506	13.	ESTS
2007	: :	A00196	He A1087	
31965		NB3629	H 93391	ES12
15120		073524	Hs 87465	Homan putative ATP/CTP-binding protein (MEAB) mRNA complete cds
28813	:	059257	Hs.91181	Human C-1 mRNA complete cds
38082	*	AA482284	H: 510483	ESTA
34723		AA287115	Hs.99697	
90	Z :	A285277	H: 8932	Morto sepiens brain expressed and finger profein mRNA complete cos
1907		24474696	H3./3043	100 FEB 44174604
1693	: :	511067	He 12544	FST.
ž.	: :	\$74728	H.7294	Andiquetin
38605	:	AA598344	H5.112492	ESTs
20967	~	N76066	Hs 35464	ESTs
24752	7.	240012	Hs 21652	Hema seplens mRNA for VIAA0567 protein complete cds
26453	:	AA621611	Hs. 70877	ESTA
ē		036076	Hs. 74763	RAN binding protein t
	. :	100000	2007	HORTO SEPRETO KNA MAN DIMONG PUBBIN CAY-LOG (CK T-KBT) FINING COMPRETE TUS
2687	: :	74/964	11154	ESTS WAY BEING A STATE FOR BOOKING GUODING PATAGO INTO SEPARATES.
1	: :	44417767		
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9	: :	143779		FST : RC 183729
78501	: :	AA137239	M. 11810	ESTA HIGHA AINDIAN BY POTHETICAL 47.4 KD PROTEIN IN SHPT. SECTT INTERGENIC REGION ISSECHARISMENT
35672	ž	AA404995		EST - RC_AAAM983
627	:	X777.48	Hs 3786	Gademate receptor metabotropic 3
26395	:	AA610064	Hs.73602	6379
36390	;	AA426291	Hs 108527	ESTS Weekly similar to No definition line found (C elegens)
21063		N93403	Hs 109443	
8	٤ :	049379	200	Human decyclycerol knase epston DGK mRNA complete cos
2 5	: :	54/1/45	10.26.01	COSTS VALSE to conference of sections of sections
26756	: :	H92977	H. 6777	Colo tropicy at the Colored Co
87.85		AA435815	H: 71965	Numer City associated RS cyclochie CARS Cyp mRNA complete cos
28.5	: =	N93185	E 55.91	ESTe
7097	:	AA911452	Hs.21840	
39462	:	D60063	He 9012	£57a
14420	:	AA600322	Hs.19574	ESTS Highly similar to AAC, RICH WRNA CLONE AACS PROTEN (Dichyostefum discoideum)
34629.	:	AA282527		EST - RC_AA282527
27.03	Z	AA429038	H1 40543	
, 6387	.	x85372	H3, 105465	
ž 5	: :	AA223874	12.14.043	Home superts makes or standing profess parties cas Templesconsesse (DNA) (Libba (1704))
	: :	4704734	4. 95773	Controlled and Controlled Control
3	: :	AA233854	H3 23348	6571
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1813 STE Modernhy univers of VY Lassessiand Index 2 [Hassians]	ES7.	551	Namen Istal Az :30-reactive done 1 (FAC1) niftNA complete cds	ES10	(S)	EST	EST - XÓJJJ?	isoleucine-RHA symbituse	Ribosomial protein (3)	ESTS MODERATED STRAFF OF THE ALIC SUBTRACTION SX YMARKING ENTRY (III) (1) SEPRENS	Marine Section of Contraction of the Contraction of Contraction of Contractions and Contraction of Contraction	aspartyc.1rma synthetase	657a	153	EST Section Sector C. N. a. D. subsects	Horns assens voltas desendent arion channel protein mRNA complete ods	H. sepiens mRNA for 8m protein G	EST s Moderaky amfar in rt/POTNETICAL 66.5 KD PROTEIN F02A9 5 IN CHROMOSOME III (Caenomabdiis alegans)		ESI 6 6 8 7 6 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8		EST	Human (GN protein mRNA complete cos	EST Weakly similar to C3088 3 (C stopana)		ESSÍS. Mente a colone anticactud comissio filosos, activated meda un lisados. Sana dominista esta	FSTs Weath and to ment of tables.	Human serine kinsse mRNA complete cts	Heapens mRNA for Workers phosphoprotein moos	Homo sepiens mRNA for MAAUSSS protein pertial ads	Signal transducer and activator of thanscription 1.alphabeta	ESTA	Figur-modelity group (modifications diversorsement) projects 2	183	Human cysteine proteese Mct2 sarform siphs (Mct2) mRNA concrete cds	ESTe			Homo suplems mRNA for ATP-dependent RNA helicase 848 complaie cds	ESTS MGHY similes to HYPOTHETICAL GTP-BUIDING PROTFIN IN PHINO-PAC2 INTERGENIC REGION (Seconaromyces carevisiae)	551	200		EST's Weakly shrifter to K0282.3 gene product (C elegans)	Human 265 protessome-essociated ped1 hamalog (POH1) mRNA complete cds	ESTs	EST. D70129	thomo Legislan suportin (mithal complete cot ESTs
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_	44279774	_		230770		HS7330			_	M6/616				-	F04370			-	_	44490969			054999	260033	M62827	A4795247	AA011134	129085	_				R08617	AA135669	AA227859	AA477045	1/808090	AA173223	NEBDS2	R10208	AA206088	207724		AA280609	AA621752	A0717N	621870	N39825
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1352	26583	7833	3674	1362	15877	19799	5946	15021	24247	40878	22323	28	87.78	41897	59586	7.0	6359	20263	14529	26767	18320	41825	4674	76661	11002	8 5	23030	41035	26895	40385	230	\$65	21241	25736	ž	2799	78877	18016	20843	1001	¥0,	4174	196.66	34.562	28448	20909	159	20340

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| 18500 | 11 | 18500 | 11 | 18500 | 1851 | 1851 | 1851 | 1851 | 1851 | 1851 | 1852 | 1851 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1852 | 1
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| 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,000,000 | 1,00
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| 17.70 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.1 | 1.
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| 150 | 11 | 150281 | 111112 | 11 | 150281 | 111112 | 11 | 111112 | 11 | 111112 | 11 | 111112 | 11 | 111112 | 11 | 111112 | 11 | 111112 | 11 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 111112 | 1111
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	7767	:	200		COTS
	34380	~	AA252414	HS.104300 ESTS	FSTs
	25101	;	R35733		EST - R35/33
	16629	~	AA036811	Hs.61859	£57¢
	25148	÷	AA028356	Hs.106106 ESTs	i EST:
	28730	~	D20859	Hs 5856	
	10200	?	R64521	Hs.77361	ESTs
	39695	Ç	AA600176	Hs 112345	. EST4
	31365	?	N67550	Hs.48907	EST
	42379	;	W37889	Hs, 100018 ESTs	ESTs
	28050	:	AA479139	Hs.75393	
	2820	;	M29474	Ha 73958	
	R927	;	AFD08442	H3.5409	
	13379	:	AA449741	Hs 4029	ESI's Weakly skralar to AF-9 PROTEIN [H.sapiens]
	52	;	U79293	Hs 90902	Human done 23948 mRNA sequence
	36575	:	AAA31085	Hs 98706	
	3	ţ	HUMTERRAM11507		AFFX-HUMFFRRM11507_5
	10970	;	AA129390	Hs 5265	EST
	25836	;	AA15230S	Hs.2248	Interfaron (gartma) induced cell line protein 10 from
	19735	;	H53038	#5.38710	EST
	40711	Ę	N53564	Hs.106159	ESTs
	4149	;	U26396	Hs 2397	RAG (recombination ectivating gene) cohort t
	5767	7	X53783	Hs.118228	
	\$500	Ş	X05232	Hs.83326	
	20310	Ę	N34893	Hs.6153	
	\$;	038145	Hs 61333	_
/	13651	ţ	W95409	Hs 59704	
	19110	•	H08778	H9.133521	
	24408	•	W80146	Hs 35962	EST _a
	26596	0,4	AA279943	Hs 88671	EST.
	32969	9	W42451	Hs \$2260	ESTS
,	27006	?	AA398695	Hs 56159	ESTa Weathy similar to E04/6 2 gare product [C alegans]
	29809	•	N21043	Hs 42932	
	9236	:	H91964	Hs 13540	
	29024	:	F09315	148,76982	Home sapiens mRNA for KIAA0583 protein partial cds
	21694	=	R39317	Hs 69403	
	37865	;	AA476623	Hs 99619	
•	1969	:	HUMTFRRM11507		AFFX+HUNTFRRM11507_3
	24862	2	241415	. Hs.6623	ESTA HEDRY GINDAR IN BONE MORPHOGENETIC PROTEIN 1 PRECURSOR (Mus musodus)
	28885	:	AA281950	Hs. 79656	ESTs .
	42300	•	135850	Hs.100703 ESTs	2 ESTs
	28	:	X92715	H# 3057	Zinc finges protein 74 (Cos52)
	39804	=	AA528803	Hs.111498 ESTs	S ESTs .
	30560	•	N49264	H\$.1334	MYB PROTO-ONCOGENE PROTEIN
	1413	:	AA600150	Hs.14366	FSTs
	38159	•	AA487021	Ms, 105703) EST
	2572	*	M27281	Hs.73793	Vescoular emodulatial growth factor
	40100	=	H75933	Hs.75901	
	20944	=	N74443	Hs 16247	· EST»
	8513	:	2446990	Ha.103135 ESTs	5 ESTs
	13877		AA475604	F3.7114	
	14509	:	AA609943	Hs 32793	
	10281	:	R80333	Hs 21182	ESTs
	25284	=	AA045074	Hs. 110146	ts. (10146 ESTs Weatly sirres to 52-kD SS-ARto automögen (H szpierty)
	6730	.	709305	_	Hs. 17154 H. septens mRNA for grotem kinase Dynk 4 partial
	18033	=	HUMISGF3A/M97935		AFFX-MUMISGF3AAM97835_MB

39242	:	AA621523	Hs.110832 ESTs	ES1s .
27.254	3.8	AA425221	Hs 81698 ESTs	ESTs
4552	3.4	U49188	Ns 76329	
18385	:	AA2272:9	Hs.110826	
16754	7.	AA046057		EST - RC_AA045067
42463	:	W60190	Hs 103135	
2 6	21	De7716	115 00315	
31795	~ :	N80703	15 50473	
22022	1,	R98192	Hs 35828	
25240	£ :	AAG39713	H3.110406	
1746	: :	038343	10079	Filmen ART-edwared phosphaticy(choinne-specific phospholipase D1a (hPLD1) mRD4 complete cds.
75921	: :	AA055552	Ne 21622	FOT What's similar to MAAAAA H sealann
3700		052201	He 1107	Heat short 10 th centain 1 (the series 10)
1604	; ;	Ī	111759	
4951	: 7		H 82321	
20418		N49209	15.32170	_
27995	9.	AA470155	Hs.75687	Homo sepiens costomer protein (COPA) mRNA complete cds
7971	•	AA287423	Hs. 126389	
27608	3.6	AA443793	H3 P4761	
24677	:	238336	H3.21201	ESTS Highly similar to POLIDVIRUS RECEPTOR HOMOLOG PRECURSOR (Mus musculus)
9328	•	D89618	Hs.3886	Homo saplens Importin-upha homolog (SRP1germns) mRNA complete cds
17678	3,0	AA134275	Hs.134510	
36209	36	AA421288	Hs.13889	ESTs Weakly similar to U.S. 1 protein [H. sepiens]
20064	ă	H98653	Hs.16056	
6179	:	144338	H3.78494	Homo sapiens mRNA for KIAA0525 protein partial cds
28622	:	011837	Hs 29846	
28628	3.8	D11888	Hs 62386	ESTa Moderately straign to PROHIBITIN [P. sepiens]
25804	3.5	AA148985	Hs.111710	ESTs
2482	\$ 6	M22898	Hs 1846	
14904	ş.	183389	Ht. 107147	_
25265	3.5	AA943765	fs 54649	
13606	3.5	AA456437	Hs,20386	
42307	2.5	196595		EST - RC_198595
¥	7.	10S068	Hs 2012	
42311	35	197257	Hs 94560	_
2023	\$2	74600	Ha.3823	INITIATION FACTOR IF-2 MITOCHONDRIAL PRECURSOR
9	7	046807	Hs 2359	Human MAP tonese phosphatase (MKP-2) mRNA complete cos
10/65	2	18287	HS 3281	Nectional postulation is
277	5.	AA063070	4000	EST - RC_AA003070_s
7437	2 :	W63/62	200	200
1300	£ ;	A4447187	1,500	the sections contribution conditions contributes a COC to a Date consistence of the contributions and contributions are contributed as the contribution of the contributions and contributions are contributed as the contribution of the contribution
3 5	: :	200	0000	restrict appears protein regulating cytokulests 1 (FYC.) Institut bompen cos
è	: :	60070	11-2000	
2 2 2	: :	20188104	2003	
1000	: :	4419074	75.27.45/	
165.10	: ;	44020128	10:1	
29,703	: :	HZ8581	Hs 97711	EST.
13838		AA465342	Hs.34045	
25585	;	AA112389	H8,107932	H(D10S170)
34018	*	AA191488	Hs.73614	
251	:	014520	Hs.64728	
3778	*	009648	Hs,383	Zinc finger protein 139 (done pHZ-37)
24535	ž	238409	Hs. 8053	ESTs
16127	:	AACOLREGO	1000	F67.

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26149 4011	::	AA250824 U20536		
986	:	X16355		NAD-DEPENDENT METHYLENETETRAHYDRJFOLATE DEHYDROGENASE
26928	:	AA342580		
36267	:	AAA24046		
32257	;	R54726	148,98493	
17365	;	AA101551	Hs 66900	
15296	2	W16694	Hs. 74284	
17675	2:	AA134064	HS.44045	
23000	7	761197	25.5	HOMO SEPONS INVINA IO KINALOZIS PETINE COMPANIE CIS
costr uzvo	: :	MA18:380	76876	
37551	: :	AA459679	136	
19.91	: 2	AA724180	H+ 67.77	
14310	: :	AASBB412	Hs.8739	
19233	:	H12634	£1.010	
12809	33	AAA24405	Hs.31839	
21555	ç	R33073	Hs.24595	ESI
13767	1,3	AA463234	Hs.119387	ES1s
4738	2	US8766	Hs.75801	
7256	:	AA075427	Ha. 17296	
17041	;	AA070364		EST - RC_AA070364
15504	2.0	W26362	Hs.44131	
18214	:	AA196635	Hs 86061	
7401	:	AA094800	Hs.55682	
18912	:	F10913	Hs.12475	
71596	61	AA425089	Hs 50722	
8410	:	H20443	Hs 31748	
2146	35	L41390 ·		EST - L41390
16683	77	F04258	Hs.38454	
33891	3.2	HUMTERRANT1507		AFFX-HUMTFRRAK11507_M
15840	3.2	X70944	Hs.91378	
3256	3.2	M92439	Hs.87157	130 KD LELXCINE-RICH PROTEIN
94	3.2	U41387	Hs.5122	Human Gu protein mRNA perial cas
21350	25	R15846	H\$ 21738	ESTs.
11981	32	AA280928	H9.24287 EST8	ESTe
23930	32	196590	Hs.125123	ESTs Weakly similar to tit! ALU SUBFAMILY J WARNING ENTRY III (H. sapiens)
30399	3.2	N45228	Hs 48495	EST
13494	22	AA453431	Hs.21043	
12908	32	AA427579	Hs 9347	
31309	7	N56818	Hs.42173	
31192	=	N54409	£ 54174	
11288	Ξ.	AA196512	Hs 25916	
9/1	5	985000	H1 62962	Intrincial to syndrase
26105	ī	AA243133	HE 46915	
11659		AA251909	Hs 36708	
19177	-	H10984	Hs 12338	£51s
17380	:	AA102568	Hs.69149	£513
42397	-	W4292B	Hs 103048 ESTs	1.551\$
14935	í	194628	Hs 63220	ESTs Weekty samiler to GA BINDING PROTEIN BETA-2 CHAIN (M septens)
3190	:	M65806	Hs.131361	
17406	ç	AA112979	Hs. 48269	Homo saptens mRNA for VRK1 complete cds
14130	:	AA489041	Hs.15140 ESTs	E51a
14134	:	A4489080	Hs 3566	Hs.3566 ESTs Highly similar to phosphorylation regulatory protein HP-10 [M.saplena]
42421	3.1	W45491	Hs 106835	
15723	2	W79060	Hs.5337	ESTs Highly similar to disosome binding protein p34 (R novegious)

2	73822	2	191715	Hs 14574	HS 14574 ESTS Highly similar to HYPOTHETICAL 103 EKD PROTEIN IN COXSB-PFK2S INTERGENIC REGION ISSACHARMINGS careastast
2	10951	3.6	AA126719	Hs 25282	
*	16891	3.6	AA292859	Ht.93567	ESIS
~	7690	**	AA215299	Hs 70930	Homb sapens chromosome 18 cosmo R30783
Ξ	1308	3.6	AA207114	HS 27842	
4	960	\$2	U24704	Hs. 111709	ts (11709 Horran antsecretory factor-1 mRNA complain cds
8	3815	7.0	AA596938		EST - RC_AA599938
Ξ	1819	3.0	AA258189	Hs 32471	
6	37433	• 2	AA454103	115,110031 EST&	ESTe
٠	2297	3.6	X13482	Hs 80506	
=	1	3.6	H59617	13 5199	ESTS FHOMY SIMILATIO UBIQUITIN-CONJUGATING ENZYME EZ-17 KD [TXOSOPNIS melanzgaster]
2	588	9	AA040382	Ms, 10290	ESTs
ñ	31574	57	N71303	Hs.50015 EST	EST
_	7614	52	AA187579	Hs 102696	15.102696 EST3 Weatdy similar to Yel0070-ap [S.cerevistree]
~	7090	20	AA005913	Hs 103300	1s 100300. Homo sapiens diptitinamide hiosynthesis protein 2 (DPH2) mRNA complote cds
~	24219	24	W69960	Hs 19418	ES1s
=	02061	۶,	H05970	Hs.133828	15.133228 Hunan done 23960 mRtA sequence
=	61771	5 2	AA136569	Hs. 144295	EST
Ξ	1362	2.8	A227251	Hs.20922	ESTs
æ	55 53	92	AA459555	Hs.31921	Homo septens mRNA for XUAA0848 protein partial cds
22	13866	• ~	AA476319	Hs.5327	ESI®
ñ	39552	26	F03605	Hs.76574	PUTATIVE 60S RIBOSOMAL PROTEIN
-	1795	56	1.13434	Hs 64162	Human chromosome 3p21.1 gene sequence complete cds
2	14740	2	050354	Hs.90315	
~ ۔۔۔	8	3.6	M64929	Hs.75200	
<u>-</u>	9962	2.6	AA430032	Hs.7487	ESTs Moderately similar to PTTG gene product (R.norvegicus)
<u>-</u>	808	52	AA171592	Hs 70980	
	2853	52	W92703	HS.103239 ESTS	ESTs
	30438	:2	N47204	Hs.46680	ESTs Weekly similar to GSOF4, 12 [C whigars]
7	5135		AA243765	Hs.76977	ESTs
÷	15457	52	w27560	Hs 90789	
~	27748	5.5	AA453159	H3 41723	
H	32315	52	R69840	Ha.70189	ESTs Weatly similar to LINE-1 REVERSE TRANSCRIPTASE HOMOLOG (Pycticabus coucang)
**	5310	3.5	AA046745	Hs.110457	
•	888	2.5	16891	Hs.79006	
¥	0.21	2.5	AA293774	Ha 21261	
m	32826	2	W20391	13 20830	
_	67 80	52	137747	Hz 59497	
7	27862	57	AA458908	Hs 50683	
-	200	57	AA065178	Hs 24382	EST
	2883	\$2	N35449	Hs.11149	
	2832	<u>.</u>	X62153	Hs. 82479	
~	2534		AA287138	Hs.59346	
•-	5157	**	U80034	Hs.68583	Human ntiochondral intermediate peptidase precursor (MIPEP) mRNA milochondral gene encoding milochondrial protein complete cds
•	ž	52	AA497013	Hs. 142592	
-	280	?	AA173505	H 35353	
-	14960	5.2	005237	Hs 99872	
	3565	\$2	AA455999	Hs.22151	
~	18185	25	AA487508	Hs 9877	
^	676	2.5	AA284744	Hs.75510	
-	7	2.5	302645	Hs. 61613	-
~	1876	\$	R43286		EST - RC_R43286
•	¥28	2.5	AA236866	Hs. 151314	I EST\$
-	60109	\$ 2	AA188381	Hs 56163	Homo sapiens rehnoblastoma-essociated protein I-EC mRNIA complete cds
•	62	≈	85026X	Hs.75914	Hs.75914. H sepiena mRNA for transmenditang projem mp24.
~	7446	5.2	AA430160	H 42765	Hs 42785 ESTs Weakly similar to F25H9.7 (C.elegens)

153	ESTs Weakly similar to WR2812.7 [C oringaris]	XYX receptor the lynosine kinase	55.18 Homo sakiens BAC ctone PG320F22 from 7671-6311-1	£51t	Human putanne ATP/GTP-binding protein (HEAB) mRNA complete cds	Human C-1 mRNA complete cds	ESTS	TOTAL SALITABLES OF ANY BASHBOOT HAY INTIGET PROPERTY MARKET CATAPOSITE CALL	E31	RAN binding protein 1	Momo septions RRM RNA binding protein Gry-rbp (GRY-RBP) mRNA complete cas	121°	Numen osteodesi stirafisting factor mRNA complete cds	ES1: NC_103729	EST a Weakly similar to No definition the found (C. elegans)	£51s	Human Citessociated RS cyclophilin CARS. Cyp mRNA complete cds	ESTs	EST\$ HIGHY SIMILATED AAC-RICH MRNA CLONE AAC3 PROTEIN (Dictyostalium discoideum)	The state of the s	THE SECRETARY TO CATE DISTRIBUTE TO THE PROPERTY OF THE PROPER	Towns deposite states of the control	ESTs	Human territinal transferase mRNA complete cds	ESTs	ESTs	Hs. 124275 Homo sapiens mRNA for KIAA0639 protein partiel cds	ESTs Highly strains to HYPOTHETICAL 40.2 KD PROTEIN K12M4.3 IN CHROMOSOME III [Ceenanabdids elegens]		רטים אינפטעל פורונים וויין אונים סטיבין אונים מיין אונים סטיבין אינונים וויין אינים סטיבין אינונים וויין אינים הראביי הראביין אינונים וויין אינים סטיבין אינונים וויין אינונים וויין אינונים וויין אינונים וויין אינונים וויי	ESTS	ESTs Weakly similar to LINEAg H-chain fusion protein (M.musculus)	ESTs Moderately shrilar to ALR [H.saptens]	ESTs Wesky similar to ORF YOR281c (S.caravisse)	ESTe Highly similar to zinc finger protein [M.muscuius]	ESTS	EDIST HEAT) HITHER TO PENCHANTE ITTE! [NOTO SEPTEMENT]	HOND SERVED AND SAIDS AND SERVED SERV	ESTA	EST	EST - RC_DS1272_s	EST	ES14	Homo Laptens mRNA for NTCF-4	ES1s	Human mRNA for RNA belicasa complate cds	100	Hs. 14811 Hamo sapiens protein lyrusine phosphalese FIR1 mRNA complete cos
		OCTA SH					10000						11,95621	73602		Hb.37747	_			1000			-			Hs.31062	Hs. 124275		900				Hs 30272	_			7CD7 54									Ht 123058		Hs 14811
N27439	AA443702	559164	N24006	F02506	U72524	059257	AA26/115	F 11087	AA621611	538076	AA253031	AA417087	183717	AA610064	AA426291	AA427745	AA435815	D80063	AA600322	AA429038	2/655A	COPOR	AA233854	W4Z788	AA459254	AA243617	R51382	AA454607	x78121	1963674	AA357394	AA343513	AA598219	AA251089	123539	N53963	N59263	A446720	AA100925	AA278907	D51272	R70012	N96254	T53138	AA309680	026528	NGORAS	W42845
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2000	27802	335	6.79	02901	15120	28813	2 2	16927	28443	452	11701	12672		28785	36390	12015	36786	39462	14420	6787	è	1497	11454	32978	27872	11623	22142	13533	i g	30.00	9763	12233	14371	26169	23065	202	307	2242	17352	11914	28799	22491	40847	42022	8053	× 4	31858	24092

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| 1472 | 1774 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 | 1772 |
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H 2102897 ESTS
H 210289 Purson univioun protein mCNA within the pSI sittion 1 carrettee cds
H 210289 Purson univioun protein mCNA within the pSI sittion 1 carrettee cds
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ESTS Weatly similar to TRANSFORMATION-SENSITIVE PROTEIN IEF SSP 3521 [H sapiens]
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HI 1511 ESTS Weekly similar to PRE-MRNA SPACING HELICASE BRR2
HI 7877 I ISHORATEST SWAND STATE STATE SHORE
HI 1877 I ISHORATEST SWAND SWAND STATE SHORE
HI 1872 EST Weekly similar to this ALU SIERARILY J WARNING SUTEN
HI 1872 EST Weekly similar to this ALU SIERARILY J WARNING SUTEN
HI 1873 EST Weekly similar to this SWAND 
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Ha.5169 ESTSV
Ha.78856 5 much
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2988 21 AV3243 H 18302 ES13
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2650 71 AV3214 H 18303 ES13
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FIGURE :

	-	Accordion	Inipene	Injune Descriptor
Frimary Key	downregulated	TORSSIGN TO	CLUSTER	
	5			
2348	֓֞֟֝֟֝֟ ֭֜֞֞	M15656	Hs.75592	Aldolase B fructose-bisphosphate
6463	>10	8060EX	Hs.74126	H.sapiens mRNA for I-15P (I-BABP) protein
42139	10	173335	Hs.93194	APOLIPOPROTEIN A-1 PRECURSOR
4544	>10	U48959	Hs.75950	H.saplens mRNA for myosin light chain kinase
1583	710	K02765	Hs.58512	COMPLEMENT C3 PRECURSOR
2426	>10	M19828	Hs.585	Apolipoprotein B (Including Ag(x) antigen)
8859	> 10	AB002351	Hs.10587	Human mRNA for KIAA0353 gene partial cds
24572	0 *0	238688	Hs.24192	ESTs
17810	01×	AA151402	Hs.46531	ESTs
2006	>10	Z70295	Hs.32966	H.saplens mRNA for GCAP-II/uroguanylin precursor
1304	×10	HG4310-		EST - HG4310-HT4580
5980	×10	X64559	Hs.65424	Tetranectin (plasminogen-binding protein)
41987	>10	T47089	Hs.121713	Cytochrome P450 subfamily XXI (steroid 21-hydroxylase congenital adrenal hyperplasia)
24461	\$10°	W94427	Hs.3807	ESTs Weakly similar to SODIUM/POTASSIUM-TRANSPORTING ATPASE GAMMA CHAIN (MUS
2372	>10	M16594	Hs.89552	Glutathione S-transferase A2
6001	91,	X65727	. Hs.89552	Glutathione S-transferase A2
1750	01.	L10955	Hs.89485	Carbonic anhydrase IV
15130	× .	U77643	Hs.95655	Homo sapiens K12 protein precursor mRNA complete cds
12467	>10	AA402656	Hs.28264	ESTs
41148	× 10	R06984		EST - RC_R06984_s
31652	۷10	N73958	Hs.50404	Human chemokine (TECK) mRNA complete cds
4605	>10	U51010		EST - U51010
28359	> 10	AA609133	Hs.58115	ESTs
24066	۰10	W32506	Hs.111676	ESTs Weakly similar to HEAT SHOCK 27 KD PROTEIN [H.Sapiens]
33282	×10	W73194	Hs.80552	ESTs
12084	. >10	AA284767	Hs.21910	ESTs
41473	01.4	R48732	Hs.11006	ESTS
27108	»10	AA404397	Hs.58414	ESTs Highly similar to ENDOTHELIAL ACTIN-BINDING PROTEIN Incinio sapiens
2848	>10	M58286	Hs.159	TUMOR NECROSIS FACTOR RECEPTOR 1 PRECURSOR
19537	×10 ×	H30270	Hs.32583	ESIS
18784	>10	F09748	Hs.7974	ESTS
5773	× 10	X54162	Hs.79386	64 KD AUTOANTIGEN D1

	•			
7/38/	>10	AA426330	HS./8264	ESIS
2866	۷10 د	M59815	Hs.76682	Complement component 4A
42530	۰10 م	W72859	Hs.74669	ESTs Weakly similar to SYNAPTOBREVIN 2 [H.sapiens]
827	۷.10 د	D87433	Hs.84753	Human mRNA for KIAA0246 gene partial cds
650	۰ د د	D78014	Hs.74566	Human mRNA for dihydropyrimidinase related protein-3 complete cds
7026	۷10 م	Z80345	Hs.127610	Acyl-Coenzyme A dehydrogenase C-2 to C-3 short chain
16736	×10	AA045306	. Hs.42996	ESTs
4630	۰10 م	U52101	Hs.9999	Human YMP mRNA complete cds
4655	۰ د د	U52969	Hs.80296	BRAIN SPECIFIC POLYPEPTIDE PEP-19
42758	۰ د د	241411	Hs.107040	ESTs
35637	¢ 13	AA402933	Hs.29283	ESTs
40392	۷. د د	H99587	Hs.108880	ESTs
7354	۰ 10	AA092348	Hs.7858	ESTs
9034	۰10	C01833	Hs.29759	ESTs Weakly similar to !!!! ALU SUBFAMILY SX WARNING ENTRY !!!! [H.sapiens]
10935	۲٠٥ ۲۰۵	AA121534	Hs.6923	ESTS Highly similar to TUBULIN BETA-5 CHAIN [Gallus gallus]
5520	01 <	X06256	Hs.119218	Integrin alpha 5 (fibronectin receptor alpha polypeptide)
2547	>10	M25809	Hs.1009	ATPase H+ transporting lysosomal (vacuolar proton pump) beta polypeptide 56/58kD isofor
9003	×10	C00808	Hs.107882	ESTs
41628	×10	R70212	Hs.79630	Immunoglobulin-associated alpha
21934	>10	R44449	Hs.48778	ESTs
11129	80	AA156873	Hs.15970	ESTs
40397	7	H00480	He 108972	TATE

FIGURE 6

Unigene Descriptor CLUSTER	Hs. 75592 Adolase B furctose-bisphosphate	Hs.74126 Hisapiens mRNA for I-15P (I-BABP) protein	HS.93194 APOLIPOPROTEIN A-I PRECURSOR	15.75950 H. sapiens mRNA for myosin light chain kinase	Hs.58512 COMPLEMENT G3 PRECURSOR	Hs.585 Apolipopratein B (including Agix) antigen)	HS.10587 Human mRNA for KIAA0353 gene partial cds	Hs.24192 ESTs	15.46531 ESTs	45.32966 H.saplens mRNA for GCAP-tl/uroguanylin precursor	_	Hs.65424 Tetranectin (plasminogen-binding protein)	Hs.121713 Cyrochrome P450 subfamily XXI (steroid 21-hydroxylase congenital adrenal hyperplasia)	Hs.3807 ESTs Weakly, similar to SODIUM/POTASSIUM-TRANSPORTING ATPASE GAMMA CHAIN [Mus musculus]	4s.89552 Glutathione S-transferase A2	Hs.89552 Ghutathione S-transferase A2	1s.89485 Carbonic anhydrase IV	Hs.95655 Homo saplens K12 protein precursor mRNA complete cds	Hs.28264 ESTs	EST - RC_R06984_s	Hs.50404 Human chemokine (TECK) mRNA complete cds	Hs. 143289 H. sapiens mRNA for metallottionein isoform 1R	EST - U51010	Hs.58115 ESTs	4s.111676 ESTs Weakly similar to HEAT SHOCK 27 KD PROTEIN [H.sapiens]	_	Hs.21910 ESTs .	_	4s.143113 Human pancreatic tipase related protein 2 (PLRP2) mRNA complete cds	Hs.37129 Sodium channel nonvoltage-gated 1 beta (Liddle syndrome)	Hs.20813 ESTs	Hs.58414 ESTs Highty similar to ENDOTHELIAL ACTIN-BINDING PROTEIN (Homo saplens)	Hs.21701 ESTs	_	Hs. 1477 Insulin-like growth factor binding protein 6	
	1		-	_	_	-					_	•	_	_	Ť	_	_	_	_	EST - RC_R06984_		_	EST - U51010		_	ш	_	_	_	••	_		_	`	_	
Accession Uniger	M15656 Hs.755	X90908 Hs.741	T73335 Hs.931	U48959 Hs.759	K02765 Hs.585	M19828 Hs.56	AB002351 Hs.105	Z38688 Hs.24*	AA151402 Hs.46	_	HG4310-	X64559 Hs.65-	T47089 Hs.121	W94427 Hs.38	M16594 Hs.89	_	L10955 Hs.89	U77643 Hs.95	AA402656 Hs.28	R06984	N73958 Hs.50	T68873 Hs.143	US1010	AA609133 Hs.58	W32506 Hs.11	W73194 Hs.80	AA284767 Hs.21	R48732 Hs.11	T29248 Hs.14	X87159 Hs.37	N64436 Hs.20	AA404397 Hs.56	AA403032 . Hs.21	T61654 Hs.93	AA079072 Hs.1	
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Human cGMP-stimulated 3'5'-cyclic nucleotide phosphodiesterase PDE2A3 (PDE2A) mRNA complete cds
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                                                                                                                                                                                                                                                                                                                                         Homo saplens cGMP-specific phosphodiesterase (PDE9A2) mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human microfibril-associated glycoprotein-2 MAGP-2 mRNA complete cds
                                                                                                                                                                                                                                                   ESTs Highly similar to ZINC FINGER PROTEIN HF.12 [Homo sapiens]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Solute carrier family 2 (facilitated glucose transporter) member 5
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Human 20-kDa myosin light chain (MLC-2) mRNA complete cds
                                                                      Endoglin (Oster-Rendu-Weber syndrame 1)
TUMOR NECROSIS FACTOR RECEPTOR 1 PRECURSOR
                                                                                                                                                                                                                                                                                                                                                                                   H.saplens KHK mRNA for ketohexokinase clone pHKHK3a
                                                   Human NF-IL-6-beta protein mRNA complete cds
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AA233369
F09748
N49281
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ESTs ESTs	ESTS	Human mRNA for EBI1-ligand chemokine complete cds	Basic fibroblast growth factor (bFGF) receptor (shorter form)	ESTs Weakly similar to SYNAPTOBREVIN 2 [H.sapiens]	Human mRNA for KIAA0246 gene partial cds	Human mRNA for dihydropyrimldinase related prolein-3 complete cds	EST	ESTs	EST	EST - RC_AA609907	ESTs	Human mRNA for KIAA0146 gene partial cds	_	N-CHIMAERIN	Matrix metalloproteinase 2 (gelatinase A 72kD gelatinase 72kD type IV collagenase)	EST - RC_R09241	ESTs	Pigment epithelium-derived factor	EST	Xanthine dehydrogenase	EST	•	_	_	ESTs Weakly similar to p20 protein [R.norvegicus]	Human mRNA for KIAA0355 gene complete cds	_	Leukotriene B4 omega hydroxylase (cytochrome P450 subfamily IVF)	SERUM AMYLOID A PROTEIN PRECURSOR	_	_	_	Human chemokine (TECK) mRNA complete cds	_	_	_		S ESTs .
Hs.17998 Hs.20733	Hs.10683	Hs.50002	Hs.748	Hs.74669	Hs.84753	Hs.74566	Hs.99289	Hs.99397	Hs.98763		Hs.17311	Hs.74670	Hs.107374	Hs.75092	Hs.111301		Hs.115726	Hs.76110	Hs.97669	Hs.250	Hs.97758	Hs.127610	Hs.118453	Hs.9693	Hs.56874	Hs.23841	Hs.108924	Hs. 101	Hs.3157	Hs.57975	Hs.42996	Hs.33455	Hs.50404	Hs.38022	Hs.13716	Hs.12112	Hs.43148	Hs.125176
AA243654 AA405199	R45577	U77180	X66945	W72859	D87433	D78014	AA452606	AA455178	AA431797	AA609907	AA487895	N57464	AA341723	R53966	AA482603	R09241	AA437388	U29953	AA399686	U39487	AA400272	280345	AA282238	N24879	AA435901	W93121	R74386	U02388	J03474	AA055163	AA045306	R85880	U86358	AA136353	AA284920	H89980	N75215	AA486185
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Hs. 101393 ESTs Moderately similar to DNAJ PROTEIN HOMOLOG 1 [Homo sapiens]	Hs.1915 PROSTATE-SPECIFIC MEMBRANE ANTIGEN	Hs.112065 EST	Hs.2048 Protease sedne 2 (trypsin 2)	Hs.76487 ESTs	Hs. 78483 Homo saplens mRNA for smoothelin	Hs.8769 ESTs	Hs. 40888 Human mRNA for KIAA0278 gene partial cds	Hs.23213 ESTs	Hs.90357 Homo sapiens teloment: repeat binding factor (TRF1) mRNA complete cds	Hs.32246 ESTs	Hs.116428 ESTs .	Hs.94413 Translent receptor potential channel 1	Hs.98998 ESTs.	Hs.8999 Human YMP mRNA complete cds	Hs.28137 ESTs	Hs.6952 ESTs .	Hs. 112961 ESTs Moderately similar to PANCREATITIS-ASSOCIATED PROTEIN 1 PRECURSOR [H.sapiens]	Hs.80296	Hs.87469 ESTs	Hs. 112125 Lecithin-cholesterol acytransferase		Hs.79368 Human epithellal membrane protein (CL-20) mRNA complete cds	Hs.107040 ESTs	Hs.29283 ESTs	_	Hs.13346 ESTs	_	Hs. 40888 Human mRNA for KIAA0278 gene partial cds	Hs.103972 Salivary proline-rich protein	Hs.26216 ESTs	Hs. 24332 ESTs Weakly similar to similar to deoxynthose-phosphate aldolase [C.elegans]	Hs.139088	Hs.34956 ESTs	Hs.25199 Home sapiens PAC done DJ130H16 from 22q12.1-qter	Hs.7858 ESTs	Hs.96744 ESTs		Hs.75860 Hydroxyacyl-Coerzyme A dehydrogenase/3-ketoacyl-Coenzyme A thiolase/encyl-Coenzyme A hydratase (trifunctional protein) alpha subunit	Hs. 106487 Homo saplens mRNA for KIAA0573 protein partial cds
AA434108	M99487	AA435805	U68061	R06986	Y13492	AA428258	D87468	H17865	U74382	AA235009	W31906	99068X	AA443311	U52101	H27852	T17215	AA399061	U52969	AA235984	R40395	AA609645	U43916	241411	AA402933	H99587	H19204	U82169	D87468	K03207	W38778	C00125	AA182845	AA283620	AA252191	AA092348	AA419011	172491	D16480	R42233
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36702	3357	36783	4876	41149	15925	12944	9317	19331	5032	34239	32852	6432	37001	4630	19489	23028	35359	4655	26030	41348	14494	4453	42758	35637	40392	19366	5184	829	1595	15574	8985	33995	7949	11670	7354	36151	42136	289	41379

-ymphotoxin-beta

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ATPase H+ transporting lysosomal (vacuolar proton pump) beta polypeptide 56/58kD (soform 1
                                                                                             ESTs Weakly similar to !!!! ALU SUBFAMILY SX WARNING ENTRY !!!! [H.sapiens]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Human done HM18 monocyte inhibitory receptor precursor mRNA complete cds
                                                                                                                                                                                                                                                                                        ESTs Weakly similar to mitogen-activated kinase kinase kinase 5 [H.sapiens]
                                                                                                                                                                                                                                                                                                                domo saplens Ca2+-dependent phospholipase A2 mRNA complete cds
                                                                                                                                                                                                                                                                ESTs Highly similar to TUBULIN BETA-5 CHAIN [Gallus gallus]
                                                                                                                                                                                                                                                                                                                                                                                 Aacrophage stimulating 1 (hepatocyte growth factor-fike)
                                                                                                                                              forno sapiens mRNA for GABA-BR1a (hGB1a) receptor
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ESTS Moderately, similar to PUTATIVE SERINE/THREONINE-PROTEIN KINASE PKWA (Thermomonospora curvata)
Laminin gamma 1 (formerly LAMB2)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human adenylyl cyclase-associated protein homolog CAP2 (CAP2) mRNA comptete cds
                   Homo sapiens kilker cell receptor (KIR103) mRNA allele ASD1 complete cds
BASIC TRANSCRIPTION FACTOR 62 KD SUBUNIT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            tuman two P-domain K+ channel TWiK-1 mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    fomo sapiens transmembrane protein mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Human msg1-related gene 1 (mrg1) mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Homo saplens Pig12 (PIG12) mRNA complete cds
                                                                                                                                                                      S100 calclum-binding protein A5 (formerly S100D)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         reacher Collins syndrome susceptibility protein
                                                                                                                               Human mRNA for KIAA0306 gene partial cds
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Homo saplens clone 23579 mRNA sequence
                                                                                                                                                                                                                                                                                                                                                                                                                                            PUTATIVE DNA BINDING PROTEIN A20
                                                                                                                                                                                                              TRANSFORMING PROTEIN RHOB
                                                                                                           Human G0S3 mRNA complete cds
                                                                                                                                                 mmunoglobulin-associated alpha
                                                                                                                                                                                                                                      Allograft inflammatory factor 1
                                                                                                                                                                                                                                                                                                                                                                                                        EST - HG2705-HT2801
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               EST - AF001359_f
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   EST - RC_R01398
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Hs.112238
Hs.7915
Hs.711233
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Hs. 111841
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AA609646
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AA058893
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T15829
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AA6021414
AA608852
U90065
AF001359
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AA479299
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MIR737	2404	×10	AA400292	Hs.23786	ESTS
10 AA460317 H5 89843 S 10 AA460317 H5 89843 S 10 AF00059 H5 10993 H 10 D12620 H5 106924 C 10 W57862 H5 106924 C 10 W57862 H5 106920 H5 106924 C 10 W69563 H5 106920 H5 58699 H5 58699 H5 58699 C 10 AA49279 H5 282813 C 10 AA438784 H5 106879 B 10 AA45847 H5 106879 B 10 AA45847 H5 106879 B 10 AA45847 H5 106879 B 10 AA45841 H5 109870 B 10 AA45848 H5 109870 B 10 AA458514 H5 109870 B 10 AA458514 H5 109870 B 10 AA458514 H5 1019 B 10 AA458514 H5 150870 B 10 AA45861 H5 180870 B 10 AA458	407	, 5	M18737	Hs.90708	GRANZYME A PRECURSOR
115445 Hs.98916 E 10 T15445 Hs.98916 E 10 AF000553 Hs.10003 H 10 D12620 Hs.106242 C 10 W67862 Hs.26941 H 10 U90543 Hs.79041 H 10 U90543 Hs.79641 H 10 AA419279 Hs.26989 H 10 AA419279 Hs.26989 Hs.79891 C 10 AA419279 Hs.26989 Hs.79891 B 10 AA4397841 Hs.106879 B 10 AA4397841 Hs.106879 Hs.7997 Hs.26991 Hs.79991 H	9820	01	AA280413	Hs.89843	Spleen focus forming virus (SFFV) proviral integration oncogene spi1
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10 AF000559 Hs.110903 H 10 D12620 Hs.106242 C 11 U80543 Hs.20841 E 11 U80543 Hs.20841 E 11 U80543 Hs.28189 E 11 U80543 Hs.28189 E 11 U805412 Hs.28189 E 12 AA419279 Hs.82813 C 13 AA45868 Hs.58889 E 14 AA45868 Hs.58889 E 15 AA419279 Hs.20819 E 16 AA45841 Hs.106879 E 17 C01384 Hs.106879 E 18 AA45841 Hs.106879 E 18 AA432381 Hs.106879 E 18 AA432381 Hs.104005 V 18 AA432381 Hs.104005 V 18 AA43338 Hs.10838 E 18 Hy.4716 Hs.11618 E 18 AA433458 Hs.109870 E 18 AA433458 Hs.109870 E 18 AA433458 Hs.1019 E 18 AA433458 Hs.1019 E 18 AA433458 Hs.1019 E 18 AA433458 Hs.1019 E 18 AA455914 Hs.1019 E 18 AA455914 Hs.1019 E 18 AA455813 Hs.50500 Hs.16500 Hs.165	1827	5	T15445	Hs.99491	H.sapiens mRNA for F2583.3 kinase like protein from C.elegans
10 D12620 Hs.106242 C Hs.106242 C Hs.21269 E Hs.22269 E Hs.22213 C Hs.21234 E Hs.2221 E Hs.22221 E Hs.22222 E Hs.2222 E Hs.22222 E Hs	82	5	AF000959	Hs.110903	Homo sapiens transmembrane protein mRNA complete cds
10 W57862 H5.2189 E	203	9	D12620	Hs.106242	CYTOCHROME P450 IVF3
10 U90543 H9.79041 H 10 LY6670 H3.109610 H 10 W80558 H8.282813 C 10 AA41927 H3.6822 H 10 AA504512 H3.6822 H 10 AA504512 H3.6829 H 10 AA504512 H3.6829 H 10 AA504512 H3.6829 H 10 AA50424 H3.106679 H 10 AA50424 H3.106679 H 10 AA50424 H3.106823 H 10 AA50424 H3.106823 H 10 AA50424 H3.10682 H 10 H1447 H3.10682 H3.104005 H 10 AA233064 H3.10839 H 11 H347 H3.10830 H 11 H347 H3.10830 H 11 H347 H3.10830 H 11 H347 H3.10987 H 11 H347 H3.10987 H 11 H347 H3.10987 H 11 H347 H3.10987 H 11 H3.1098 H 11 H3.109 H 11 H3.1098 H 11 H3.1098 H 11 H3.1098 H 11 H3.1098 H 11 H3.109	4159	5	W57862	Hs.21289	ESTs
10	302	5	U90543	Hs.79041	Human butyrophilin (BTF1) mRNA complete cds
10	219	õ	L76670	Hs.109610	Homo sapiens nkat7 mRNA complete cds
10 AA419279 Hs.82813 C 10 AA45866 Hs.95898 E 9 F10660 Hs.103679 E 10 AA45841 Hs.106879 E 10 AA45841 Hs.106879 E 10 AA45841 Hs.106879 E 10 AA45841 Hs.106870 E 10 CO1384 Hs.106870 E 10 CO1384 Hs.106870 E 10 CO1384 Hs.106870 E 10 AA432381 Hs.97357 E 10 AA433281 Hs.10689 E 10 AA433257 Hs.1069870 E 10 AA433257 Hs.25511 Hs.1079 E 10 AA453458 Hs.1079 E 10 AA453458 Hs.1079 E 10 AA453614 Hs.1079 E 10 AA453614 Hs.1079 E 10 AA453614 Hs.5063 Hs.25060 Hs.25060 Hs.25060 Hs.25060 Hs.25060 Hs.25060 Hs.26060	4392	₽	W88568	Hs.58589	Homo saplens glycogenin-2 gamma (glycogenin-2) mRNA complete cds
10 AA504512 Hs,76852 Hs,76852 Hs,76852 Hs,96898 Hs,96898 Hs,96898 Hs,96898 Hs,96898 Hs,96898 Hs,96898 Hs,96898 Hs,96898 Hs,96899 Hs,96899 Hs,96899 Hs,96899 Hs,96999 Hs,9699 Hs,96999 Hs,96999 Hs,96999 Hs,96999 Hs,9699 Hs,96999 Hs,96999 Hs,96999 Hs,9699 Hs,969	6159	5	AA419279	Hs.82813	Colony-stimulating factor 1 (M-CSF)
10 AA458666 Hs,95898 E F10640 Hs,12354 E A337841 Hs,106879 E A4101632 Hs,22971 E A4537841 Hs,106530 Hs,22971 E E E E E E E E E E E E E E E E E E E	8251	5	AA504512	Hs.76852	ESTs Weakly similar to ZK792.1 [C.elegans]
F10640	7592	9	AA458668	Hs.95898	ESTs
AA397841 Hs.106679 E AA4101632 Hs.22971 B AA455474 Hs.100530 B WYG2272 Hs.25601 Hs.106823 Hs.106823 Hs.106823 Hs.106823 Hs.106823 Hs.106823 Hs.106823 Hs.106823 Hs.25601 Hs.106829 Hs.10683 Hs.25601 Hs.10683 Hs.25601 Hs.10683 Hs.25601 Hs.10683 Hs.25601 Hs.10683 Hs.25601 Hs.10683 Hs.25601 Hs.25601 Hs.25601 Hs.25601 Hs.25601 Hs.25601 Hs.26603 Hs.25603 Hs.25600 Hs.25600 Hs.26603 Hs.25600 Hs.26603 Hs.26600 Hs.2	9619	ø	F10640	Hs.12354	ESTs
8 AA101632 Hs, 22971 B AA452474 Hs, 100530 B W92272 Hs, 25601 Hs, 100530 B W92272 Hs, 25601 Hs, 100822 Hs, 100822 Hs, 100822 Hs, 100822 Hs, 100930 B Hs, 1009300 B Hs, 100930 B Hs, 1009300 B Hs,	3240	on	AA397841	Hs.106679	EŚTs
MA455474 Hs. 100530	0887	æ	AA101632	Hs.22971	ESTs
9 W92272 Hs.25601 H 9 S77763 Hs.106823 H 9 S77763 Hs.106823 H 9 T56281 Hs.10400 H 156281 Hs.110440 H 9 H1476 Hs.110440 H 9 H1476 Hs.11615 H 10832 Hs.10839 H 10832 Hs.10839 H 10832 Hs.25511 H 10832 Hs.25500 H 10832 Hs.25050 H 10832 Hs.25050 H 10832 Hs.25050 H 10832 Hs.25050 H 10832 Hs.25050 H 10832 Hs.25050 H	7500	d i	AA455474	Hs.100530	ESTs ·
9 S77763 Hs.106823 H 9 AA432381 Hs.104005 H 9 T56281 Hs.104005 H 9 AA23282 Hs.10400 H 1 T56281 Hs.10400 H 9 AA23064 Hs.10839 H 1 Hs.10440 Hs.10839 Hs.10830 H 1 MA2733 Hs.10830 Hs.25511 Hs.10830 Hs.25511 Hs.25011 Hs.25511 Hs.25011 Hs.25511 Hs.25511 Hs.25511 Hs.25511 Hs.25511 Hs.25511 Hs.25011 Hs.25511	2650	Ø	W92272	Hs.25601	Homo sapiens zinc-finger helicase (hZFH) mRNA complete cds
9 S77763 Hs,75643 H 9 SA433381 Hs,9737 F 9 T56281 Hs,104005 N 9 AA259064 Hs,10839 H 17476 Hs,11615 H 17476 Hs,11615 H 17476 Hs,11615 H 17476 Hs,11615 H 17476 Hs,10830 H 17476 Hs,25511 H 17476 Hs,25051 H	9011	69	C01394	Hs.106823	Homo sapiens done 24818 mRNA sequence
9 AA432381 Hs.97357 ES 75892 Hs.104005 VS.76892 Hs.10440	3490	o	S77763	Hs.75643	Homo sapiens NF-£2 protein (NF-£2) mRNA complete cds
9 \$78992 Hs.10400 V 9 \$A25906 Hs.10440 Hs.10939 H17476 Hs.11615 Hs.10939 Hs.10939 Hs.10939 Hs.10930 Hs.25511 Hs.25511 Hs.25511 Hs.25511 Hs.25511 Hs.25511 Hs.25514 Hs.1019 Hs.25514 Hs.1019 Hs.25514 Hs.2501 Hs.25514 Hs.2501	6691	Œ	AA432381	Hs.97357	ESTs Moderately similar to NITRIC OXIDE SYNTHASE INDUCIBLE [H.sapiens]
156281 H5.110440 H5.25061 H5.110440 H5.25064 H5.10839 H5.25511 H5.	3478	69	S76992	Hs.104005	Vav 2 oncogene
AA259064 Hs. 10839 E H17476 Hs. 11615 B W42733 Hs. 10830 E B WA233257 Hs. 25511 H B W431337 Hs. 28017 E AA431337 Hs. 28017 E AA455914 Hs. 1019 E X65644 Hs. 7301 E X55448 Hs. 7301 E X5548 Hs. 7301 E	2034	Ch.	T56281	Hs.110440	Human metallothionein (MT)I-F gene
H17476 H5.11615 W42733 H5.10870 W42733 H5.10870 W19008 H5.75511 A4433257 H5.25511 A4453458 H5.7301 A4455914 H5.1019 X5544 H5.75053 W93074 H5.5053 K5544 H5.75063 W3544 H5.75063	1845	æ	AA259064	Hs. 10839	ESTs Weakly similar to unknown (S.cerevistae)
M42733 Hs.109870 M423257 Hs.25511 M433257 Hs.25511 M4431337 Hs.98017 M4455914 Hs.1019 M55644 Hs.7301 M93074 Hs.59342 M93074 Hs.59342 M45648 Hs.718 M5648 Hs.80500 M5648 Hs.80500	9317	cts	H17476	Hs.11615	ESTs Weakly similar to dual specificity phosphatase [H.sapiens]
8 AA233257 Hs.25511 Hs.26511 H	2395	•0	W42733	Hs.109870	ĖSTs
8 AA431337 Hs.98017 B AA453458 Hs.7301 B AA455914 Hs.7301 B AA455914 Hs.7301 B AA55944 Hs.7301 B AA55848 Hs.3118 Hs.55060 B N21684 Hs.58060 B N21684 Hs.80500 Hs.8050	1425	œ	AA233257	Hs.25511	Homo saplens mRNA for HIc-5 partial cds
8 AA431337 Hs.98017 R 8 AA453458 Hs.7301 B 8 AA455914 Hs.1019 R 8 W93074 Hs.53942 B 8 X55448 Hs.3118 B AA156873 Hs.15970 B N94551 Hs.53060 B N21684 Hs.80500 B	5310	80	W19098	Hs.7921	ESTs
AA453458 Hs.7301 8 AA455914 Hs.1019 8 X65644 Hs.7063 4 W93074 Hs.59042 B X55448 Hs.3118 Hs.8916 B AA156873 Hs.59060 B N21684 Hs.80500 Hs.8	10991	6 0	AA431337	Hs.98017	ESTs
AA455914 Hs.1019 RS.65644 Hs.75063 Hs.75063 Hs.75048 Hs.73042 Hs.7318	3499	6 0	AA453458	Hs.7301	ESTs
8 X65644 Hs.75063 H 8 W93074 Hs.59342 E 8 X55448 Hs.3118 H 8 AA156873 Hs.16970 B 9 N94551 Hs.55060 H 9 N21684 Hs.80500 H	17514	e 0	AA455914	Hs.1019	Parathyroid hormone receptor 1
8 W93074 Hs.59342 6 8 X55448 Hs.3118 1 8 AA156873 Hs.15970 1 8 N94551 Hs.55060 1 9 N21684 Hs.80500 1	5998	80	X65644	Hs.75063	HUMAN IMMUNODEFICIENCY VIRUS TYPE I ENHANCER-BINDING PROTEIN 2
8 X55448 Hs.3118 I 8 AA156873 Hs.15970 I 8 N94551 Hs.55060 I 9 N21684 Hs.80500 I	3589	100	W93074	Hs.59342	ESTs
8 AA156873 Hs.15970 I 8 N94551 Hs.55060 I 8 N21684 Hs.80500 I	5801	€0	X55448	Hs.3118	H.sapiens mRNA for 2.19 gene
8 N94551 Hs.55060 (1129	8 0	AA156873	Hs.15970	ESTs
8 N21684 Hs.80500	1987	s 0	N94551	Hs.55060	ESTs
	0438	6 0	N21684	Hs.80500	Human mRNA for KIAA0061 gene partial cds
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Homo sapiens cone 23904 mRNA sequence Sorting nexin 1 Human clone IMAGE:35527 unknown protein mRNA partial cds Human clone IMAGE:35527 unknown protein partial cds Human clone IMAGE:35527 unknown protein partial cds ESTs STATHMIN ESTs Human clone IMAGE:35527 unknown protein partial cds ESTs STATHMIN ESTs Human clone IMAGE:35527 unknown protein partial cds ESTs Surfactant pulmonary-associated protein D Propertir P factor complement Human light-activated protein IMAs sequence Cabindan 2 (29kD calretirin) ESTs ESTs ESTs ESTs Human mRNA for ROX protein ESTs Human light-activated protein kinase PRK1 mRNA complete cds Glycogen synthase 1 (muscle) Thromboxane A2 receptor ESTs ESTs Human mRNA for KIAA0385 gene complete cds ESTs ESTs Human mRNA for KIAA0385 gene complete cds ESTs ESTs ESTs FSTs FSTs Human mRNA for KIAA0385 gene complete cds ESTs ESTs ESTs ESTs ESTs FSTs FSTs FSTs FSTs FSTs FSTs FSTs F	Hs. 75283 Hs. 87197 Hs. 7456 Hs. 199410 Hs. 106385 Hs. 106385 Hs. 106371 Hs. 27373 Hs. 27973 Hs. 19610 Hs. 19610 Hs. 19638 Hs.	AAA28839 C01380 U85922 AA284403 AA46114 D66419 AA466112 AA262308 AA460127 T40652 D45608 M83622 H24466 AA425782 W66410 T86828 AA486319 AA460319 AA401052 U33053 T15833 N21819 D3601 AA401052 T33137 AA46014 AA40162 AA40162 T33137 AA26144 AA46014 AA40162 T33137 AA26144 AA40162 AA40162 AA40162 T33137 AA26164 AA40162		11599 90010 90010 90010 17953 17953 170617 19529 11686 37294 223201 22196 12376 12376 12376 12376 12376 12376 12376 12376 12376 12376 12174 1517 1517 1517 1517 1517 1517 1517 1
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_	Hs.19167	N69989	4	863
- 1	HS.9500	ASSEUR	•	4
-	10.050	000000	•	
_	Hs.10435	AA251547	•	98
	Hs.7967	T33137	•	920
_	Hs.11859	AA490182	*	28
_	HS. HOS. S.	AA349990	4	257
- L	13.0300	100001	•	z ;
•	70000	100000	•	
Ĭ	Hs.772	J04501	4	<u>-</u>
-	Hs.14896	H21819	4	क्र
	Hs.30732	AA410529	•	9
_	Hs.25497	N21380	4	Ξ
_	Hs.100227	T15833	•	4
_	Hs.2499	U33053	4	8
•		AFFX-	4	8
ш	Hs.48469	AA460319	4	94
٠.	Hs.19610	AA399271	4	92
	Hs.108169	180628	4	7
_	Hs. 106857	W68410	₹	88
-	Hs.27973	AA425782	4	8
_	Hs.85053	H24456	₹	8
	Hs.53155	M83652	4	2
••	HS.83792	D45608	'n	2
	HS.82/9	140652	۰ م	ا چ
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_	Hs 110571	AA450127	ur.	ő
_	Hs.106385	AA262308	v n	88
_	Hs.99410	AA456112	10	53
	Hs.81915	D60419	S	8
_	Hs.55409	AA446114	ĸ	14
_	Hs.74750	AA284403	w	ß
_	Hs.87197	U85992	v	\$
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13522	4	AA454115	Hs.6000	ESTs
18444	4	AA232646	Hs.68061	ESTs
27665	٣	AA447759	Hs.134724	ESTs'
21382	33	R16896	Hs.89615	PROTEIN KINASE C THETA TYPE
2052	m	L36818	Hs.75339	Inositol polyphosphate phosphatase-like protein 1 (51C protein)
9039	m	C02049	Hs.106291	ESTs
34888	ന	AA303078	Hs.94479	Human GT334 protein (GT334) gene mRNA complete cds
11047		AA142849	Hs.22660	ESTs
19451	ب	H23747	Hs.31697	ESTs
2822	6	M55621	Hs.117946	N-acetylglucosaminyttransferase I ·
13928	ဗ	AA478441	Hs.11590	ESTs
12064	6	AA283848	Hs.11367	ESTs Weakly similar to KIAA0009 [H.sapiens]
3836	က	U12707	Hs.2157	Wiskott-Aldrich syndrome (ecezema-thrombocytopenia)
4528	m	U48251	Hs.75871	Homo sapiens protein kinase C-binding protein RACK7 mRNA partial cds
42064	က	T63364	Hs.9225	ESTs Highly similar to FIBRINOGEN-LIKE PROTEIN A PRECURSOR [Parastichopus parvimensis]
4596	es	U50743	Hs.19520	Sodium/potassium ATPase gamma subunit
4914	က	U67611		EST - U67611
20168	6	N24106	Hs.2799	Cartilage linking protein 1
24281	e	W79773	Hs.16511	ESTs
19634	က	H44866	Hs.31597	ESTs
10989	င	AA132366	Hs.8023	Homo sapiens mRNA for SPOP
6587	e	X97748		EST - X97748
14096	က	AA487558	Hs.8135	ESTs
13350	m	AA449297	Hs.8944	ESTs Weakly similar to type 1 procollagen C-proteinase enhancer protein precursor [H.sapiens]
33930	က	AA169539	Hs.95870	ESTs
34215	m	AA233855	Hs. 104252	UTROPHIN
22509	٣	R71393	Hs.29190	ESTs
20065	ო	H98657	Hs.27291	ESTS
31091	n	N63076	Hs.138746	_
2493	က	M22919	Hs.77385	MYOSIN LIGHT CHAIN ALKALI SMOOTH-MUSCLE ISOFORM
28913	ო	F01560	Hs.22583	ESTs Highly similar to co-repressor protein (M.musculus)
14323	က	AA598575	Hs.12851	ESTs
34914	8	AA338729	Hs.133096	
14236	က	AA496891	Hs.5011	ESTs Weakly similar to The ha1237 gene product is related to S.pombe rad21 gene product. [H.sapiens
24594	က	Z38804	Hs.22555	ESTs Highly similar to OPIOID BINDING PROTEIN/CELL ADHESION MOLECULE PRECURSOR (Box
22589	ო	R79580	Hs.29874	ESTs
22156	m	R52145	Hs.25894	ESTs
16404	e	AA021284	Hs.60554	ESTs Moderately similar to sodium-calcium exchanger form 3 [R.norvegicus]
29310	es	H66642	Hs.88729	ESTs

Acyl-Coenzyme A dehydrogenase C-2 to C-3 short chain

Probable transcription factor PML (alternative products)

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-tuman clone ID 193225 NAD (H)-specific isocitrate dehydrogenase gamma subunit mRNA alternatively spliced partial cds
                                                                                 Human LIM protein MLP mRNA complete cds
IMMUNOGLOBULIN-RELATED 14.1 PROTEIN PRECURSOR
                                                                                                                                                                                                                                                  STs Weakly similar to No definition line found [C.elegans]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ESTs Weakly similar to hypothetical protein [H.sapiens]
                                                                                                                                                                                                                                                                                                                                                           Homo saplens bicaudat-D (BICD) mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Transcription factor COUP 2 (a.k.a. ARP1)
EXTRACELLULAR SIGNAL-REGULATED KINASE 3
                                                                                                                                                                                                                                                                                                                                                                                                                   Phosphodiesterase 6A cGMP-specific rod alpha
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Homo sapiens done 24440 mRNA sequence
Homo sapiens done 24525 mRNA sequence
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                                                                                                                                       Vitric oxide synthase 3 (endothelial cell)
                                                                                                                                                                                                                                                                            CDW52 antigen (CAMPATH-1 antigen)
                                                                                                                                                                                                                        Protein kinase C substrate 80K-H
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ST - HG1804-HT1829
                                                                                                                                                                                                                                                                                                                                      EST - RC_AA063316
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ESTs	DNA-BINDING PROTEIN A	ESTs	ESTs	Peroxisornal biogenesis factor 6	Human RGP3 mRNA complete cds	ESTs	ESTs	ESTs	Homo sapiens germline mRNA sequence	ESTs	ESTs	ESTs	EST - HG3227-HT3404	EST - RC_AA401489	H.sapiens HD21 mRNA	ESTs.	Human mRNA for TPRD complete cds	KERATIN TYPE II CYTOSKELETAL 6D	ESTs	ESTs	ESTs	Human phospholipase c delta 1 mRNA complete cds	ESTs	EST - AA428531	EST - RC_AA128926	ESTs	Homo saplens ezrin-radixin-moesin binding phosphoprotein-50 mRNA complete cds	ESTs	ESTs	N-ACETYLLACTOSAMINE SYNTHASE	Homo sapiens vesicle trafficking protein sec22b mRNA complete cds	ESTs	ESTs	ESTs	Human peroxisome proliferator activated receptor mRNA complete cds	ESTS	Jun D proto-oncogene	AFFX-HSAC07/X00351_M	ESTs Weakly similar to F35G12.9 [C.elegans]
Hs.24812	Hs.89491	Hs.77978	Hs. 19978	Hs.30729	Hs.82294	Hs.5723	Hs.7934	Hs. 100530	Hs.12840	Hs.22222	Hs.25046	Hs.22410			Hs.137591	Hs.26812	Hs.75395	Hs.111758	Hs.112751	Hs. 104965	Hs.57475	Hs.80776	Hs.32699			Hs.121515	Hs.3354	Hs.109727	Hs.11809	Hs.80881	Hs.50785	Hs.25604	Hs.36574	Hs.64001	Hs.106415	Hs.65311	Hs.2780		Hs.34769
Z40923	X95325	AA402495	N52322	D83703	U27655	AA449716	AA480045	R59906	T33164	AA213667	Z38888	AA437225	HG3227-	AA401489	Z49105	N59373	AA007509	L42611	AA609707	AA478162	D81123	U09117	H37834	AA428531	AA128926	H18829	AA174185	AA059099	AA490620	H85120	R64199	H27675	AA029703	N53143	L07592	AA411473	AA11550B	AFFX-	AA125969
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24819	6532	27085	20487	724	4132	13375	13988	22306	23167	11320	24608	13163	1139	35572	6964	30963	16164	2174	38958	37919	28905	3745	19545	8416	17569	19354	7598	25385	14176	29487	10197	19488	10568	30799	9638	27195	17438	24932	10944

ESTS	ESTs	ESTs	Homo sapiens mRNA for tyrosyf sulfotransferase-2	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs Highly similar to RIBOSOMAL PROTEIN S6 KINASE [Homo saplens]	ESTs Weakly similar to HYPOTHETICAL PROTEIN KIAA0168 [H.sapiens]	AFFX-HUMGAPDH/M33197_M	H.sapiens mRNA for chloride channel (putative) 2139bp	ESTs Weakly similar to zinc finger protein [H.sapiens]	ESTs	EST - RC_H82929	ESTs	ESTs:	ESTs	ESTs	ESTs	ESTs Weakly similar to Lph17p [S.cerevisiae]	ESTs	Homo sapiens mesoderm-specific basic-helix-loop-helix protein (POD1) mRNA complete cds	EST - RC_AA435753	ESTs	ESTs	EST - RC_AA129856	ESTs Highly similar to THREONYL-TRNA SYNTHETASE CYTOPLASMIC [Homo sapiens]	ESTs	Homo sapiens mRNA for NA14 protein	ESTs	ESTs	Cholinergic receptor nicotinic detta polypeptide	ESTs	ESTs Weakly similar to MITOTIC MAD2 PROTEIN [S.cerewisiae]	Human mRNA for KIAA0321 gene partial cds	ESTs	ESTs Highly similar to c-Jun leucine zipper interactive [M.musculus]	EST
Hs.142702	Hs.8245	Hs.24545	Hs.26350	Hs.16466	Hs.79788	Hs.54865	Hs.71626	Hs.21782	Hs.103081	Hs.32822		Hs.123123	Hs. 133475	Hs.31562		Hs.6217	Hs.97602	Hs.102755	Hs.56782	Hs.91202	Hs.27262	Hs.35096	Hs.78061		Hs.110783	Hs.20573		Hs.107365	Hs.11759	Hs.18528	Hs.15548	Hs.20102	Hs.99975	Hs.77480	Hs.19400	Hs.8663	Hs.104476	Hs.10552	Hs.97682
T98199	AA287665	AA421050	AA459389	AA430474	AA094921	AA404707	AA135941	F04686	AA410355	AA291786	AFFX-	230643	AA402267	H46074	H82929	F0444	AA398161	R53520	D59267	AA151480	Z39191	W51743	W73859	AA435753	R97176	AA069425	AA129856	N32118	AA609045	AA426521	H53059	AA609346	X55019	AA232508	R54743	AB002319	AA477891	AA402493	AA399593
8	7	7	7	7	7	8	7	~	. 7	7	8	-	-	_	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-			-	-	-	-	-		-	-
42324	34756	12743	13676	13009	7403	35669	17701	18713	8314	7990	42791	6893	35607	9468	29469	18692	35205	22184	28815	17813	24655	15611	15700	36770	32400	10802	17593	20266	14447	12892	19738	14471	5796	18441	10164	8830	8682	35620	35401

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ESTs Weakly similar to HYPOTHETICAL 88.1 KD PROTEIN K02D10.1 IN CHROMOSOME III [C.elegans]
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Homo sapiens p38beta2 MAP kinase mRNA complete cds ESTs ESTs
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THY-1 MEMBRANE GLYCOPROTEIN PRECURSOR
ESTs Weakly similar to No definition line found {C.elegans}
                                                                                                                                                                                              ESTs Weakly similar to No definition line found [C.elegans]
Homo sapiens clone 24800 mRNA sequence
Homeo box B5 (2.1 protein)
                                                                                                                                                                                                                                                                                    Homo sapiens done 23565 unknown mRNA partial cds
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                                                                                                                                          Human mRNA for KIAA0296 gene complete cds
                                                                                                                  Ribosomal protein S28
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Hs. 7985
Hs. 12600
Hs. 44608
Hs. 107725
Hs. 107725
Hs. 2624
Hs. 2624
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Hs. 29126
Hs. 8961
Hs. 8961
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Hs. 14593
Hs. 84344
Hs. 87225
Hs. 110095
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Hs. 6448
Hs. 55443
Hs. 25443
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Hs.34183
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T96407 Hs.17812 ESTs

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FIGURE 7

Unigene Descriptor	Aldolase B fructose-bisphosphate	H.saplens mRNA for t-15P (I-BABP) protein	APOLIPOPROTEIN A-I PRECURSOR	H.sapiens mRNA for myosin light chain kinase	COMPLEMENT C3 PRECURSOR	Apolipoprotein B (including Ag(x) antigen)	Human mRNA for KIAA0353 gene partial ods	ESTs	ESTs	H.sapiens mRNA for GCAP-tifuroguanylin precursor	EST - HG4310-HT4580	Tetranectin (plasminogen-binding protein)	Cytochrome P450 subfamily XXI (sterold 21-hydroxylase congenital adrenal hyperplasia)	ESTs Weakly simitar to SODIUM/POTASSIUM-TRANSPORTING ATPASE GAMMA CHAIN [Mus musculus]	Glutathione S-transferase A2	Glutathione S-transferase A2	Carbonic anhydrase IV	Homo saplens K12 protein precursor mRNA complete cds	ESTs	EST - RC_R06984_s	Human chemokine (TECK) mRNA complete cds	H saplens mRNA for metallothionein isoform 1R	EST - U51010	ESTs	ESTs Weakly similar to HEAT SHOCK 27 KD PROTEIN [H.sapiens]	ESTs	ESTs	ESIs ·	Human pancreatic lipase related protein 2 (PLRP2) mRNA complete cds	Sodium channel nonvoltage-gated 1 beta (Liddle syndrome)	ESTs	ESTs Highly similar to ENDOTHELIAL ACTIN-BINDING PROTEIN [Homo sapiens]	EST\$	APOLIPOPROTEIN A-I PRECURSOR	Insulin-IIko growth factor binding protein 6	ESTS
Unigene CLUSTER	Hs.75592	Hs.74126	Hs.93194	Hs.75950	Hs.58512	Hs.585	Hs.10587	Hs.24192	Hs.46531	Hs.32966		Hs.65424	Hs.121713	Hs.3807	Hs.89552	Hs.89552	Hs.89485	Hs.95655	Hs.28264		Hs.50404	Hs.143289		Hs.58115	Hs.111676	Hs.80552	Hs.21910	Hs.11006	Hs.143113	Hs.37129	Hs.20813	Hs.58414	Hs.21701	Hs.93194	Hs.1477	Hs.78293
Accession	M15656	X90908	T73335	U48959	K02765	M19828	AB002351	. 238688	AA151402	270295	HG4310-	X64559	T47089	W94427	M16594	X65727	L10955	U77643	AA402656	R06984	N73958	168873	U51010	AA609133	W32506	W73194	AA284767	R48732	T29248	X87159	N64436	AA404397	AA403032	T61654	AA079072	AA303081
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Human cGMP-stimulated 3'5'-cyclic nucleotide phosphodiesterase PDE2A3 (PDE2A) mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human apM1 mRNA for GS3109 (novel adipose specific collagen-like factor) complete cds
                                                                                                                                                                                                                                                                                                                                                                                              Homo saplens cGMP-specific phosphodiesterase (PDE9A2) mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Human microfibril-associated glycoprotein-2 MAGP-2 mRNA complete cds
                                                                                                                                                                                                                                                                                          ESTs Highly similar to ZINC FINGER PROTEIN HF.12 [Homo sapiens]
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                                                                                                      TUMOR NECROSIS FACTOR RECEPTOR 1 PRECURSOR
                                                                                                                                                                                                                                                                                                                                                                                                                                               H.sapiens KHK mRNA for ketohexokinase done pHKHK3a
                                                     Human NF-IL6-beta protein mRNA complete cds
                                                                             Endoglin (Osler-Rendu-Weber syndrame 1)
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>10 U02388 Hs.101 >10 F08876 Hs.8008 >10 R73075 Hs.29327 >10 AA45580 Hs.39405 >10 R03483 Hs.13850 >10 U42031 Hs.7557 >10 M14777 Hs.88552		Hs.108924	R74386	×10	47
>10 R74386 Hs. 108924 >10 U02388 Hs. 101 >10 F08876 Hs. 8008 >10 R73075 Hs. 29327 >10 AA455960 Hs. 93405 >10 R05483 Hs. 138500 >10 U42031 Hs. 7557 >10 M14777 Hs. 88552		Hs.23841	W93121	×10	47
>10 W93121 Hs.23841 >10 R74386 Hs.106924 >10 R74386 Hs.101 >10 F08876 Hs.8008 >10 R73075 Hs.29327 >10 AA455960 Hs.39405 >10 R05483 Hs.138500 >10 U42031 Hs.7557 >10 M14777 Hs.88552		Hs.56874	AA435901	>10	છ
>10 AA435901 Hs.56874 >10 W93121 Hs.23841 >10 W93121 Hs.20841 >10 U62388 Hs.101 >10 F08876 Hs.8008 >10 R73075 Hs.29327 >10 AA455860 Hs.99405 >10 AA455861 Hs.338500 >10 U42031 Hs.7557 >10 M14777 Hs.88552	ESTs	Hs.9693	N24879	×10	6
>10 NA4879 Hs.9893 >10 AA435901 Hs.58874 >10 W93121 Hs.23841 >10 R74386 Hs.10924 >10 R74386 Hs.10924 >10 F08876 Hs.8008 >10 R73075 Hs.28008 >10 AA455960 Hs.99405 >10 R05483 Hs.13580 >10 R02031 Hs.7557 >10 M14777 Hs.88552		Hs.118463	AA282238	×10	بي
>10 AA28238 Hs. 118463 >10 N24879 Hs. 5683 >10 AA435901 Hs. 56874 >10 W93121 Hs. 123841 >10 R74386 Hs. 106924 >10 U02388 Hs. 101 >10 F08876 Hs. 2008 >10 R73075 Hs. 2008 >10 AA455960 Hs. 30405 >10 R05483 Hs. 133850 >10 U42031 Hs. 7557 >10 W14777 Hs. 88552	_	Hs.127610	Z80345	×10	9
>10 Z80345 Hs. 127610 >10 AA28228 Hs. 118463 >10 AA435901 Hs. 56874 >10 AA435901 Hs. 56874 >10 W93121 Hs. 10924 >10 R74386 Hs. 10924 >10 U02388 Hs. 101 >10 F08876 Hs. 2008 >10 AA45596 Hs. 29327 >10 AA45596 Hs. 33850 >10 R03483 Hs. 138500 >10 U42031 Hs. 7557 >10 M14777 Hs. 88552		Hs.97758	AA400272	>10	ន
>10 AA400272 Hs.97758 >10 A280345 Hs.127610 >10 A282238 Hs.11863 >10 NZ4879 Hs.58874 >10 AA435901 Hs.58874 >10 AA435901 Hs.23841 >10 W93121 Hs.23841 >10 R74386 Hs.10924 >10 U02388 Hs.101 >10 R73078 Hs.29327 >10 AA455960 Hs.939405 >10 AA455960 Hs.33850 >10 R03463 Hs.13850 >10 HA777 Hs.88552	Xanthine dehydrogenase	Hs.250	U39487	•>10	80
->10 U39487 H-S.250 ->10 AA400222 H-S.97758 ->10 AA28223 H-S.118463 ->10 N24879 H-S.9693 ->10 N24879 H-S.9693 ->10 AA435901 H-S.6874 ->10 W93121 H-S.2341 ->10 R74386 H-S.10824 ->10 R74386 H-S.101 ->10 R74386 H-S.101 ->10 R74386 H-S.10387 ->10 R74386 H-S.99327 ->10 R74386 H-S.99327 ->10 R744386 H-S.99327 ->10 R744386 H-S.99405 ->10 R744386 H-S.99405 ->10 AA455960 H-S.99405 ->10 R744386 H-S.99405 ->10 R744586 H-S.99405 ->10 R744587 H-S.99405		Hs.97669	AA399686	×10	Ξ.
>10 AA399686 Hs.97669 >10 AA402021 Hs.9768 >10 AA40221 Hs.13461 >10 AA282238 Hs.118463 >10 NZ4879 Hs.9693 >10 NZ4879 Hs.58874 >10 NZ43501 Hs.58874 >10 NZ43501 Hs.20924 >10 R74366 Hs.10924 >10 R74366 Hs.10924 >10 R74366 Hs.8008 >10 R73075 Hs.2037 >10 R73075 Hs.30405 >10 R74559 Hs.13550 >10 R745361 Hs.13550 >10 R745360 Hs.94552 >10 R74777 Hs.88552		Hs.76110	U29953	>10	2
>10 U29953 Hs.76110 >10 AA39666 Hs.97669 >10 AA400212 Hs.97669 >10 AA400212 Hs.97758 >10 AA400212 Hs.127610 >10 AA282238 Hs.118463 >10 N24879 Hs.9683 >10 N24879 Hs.9683 >10 W43121 Hs.23841 >10 W43121 Hs.20814 >10 W43121 Hs.20814 >10 W43121 Hs.20814 >10 R74386 Hs.10624 >10 R74386 Hs.10624 >10 R74386 Hs.30940 >10 R74556 Hs.39405 >10 R745366 Hs.7557 >10 R745366 Hs.7557 >10 R745366 Hs.38552		Hs.115726	AA437388	>10	12
>10 AA437388 Hs.11572 >10 U29953 Hs.76110 >10 U39886 Hs.9768 >10 U39487 Hs.250 >10 AA400272 Hs.13768 >10 AA400272 Hs.137610 >10 AA282238 Hs.118463 >10 N24879 Hs.56874 >10 N4345901 Hs.56874 >10 W431501 Hs.103841 >10 U02386 Hs.106924 >10 P68876 Hs.8068 >10 R73075 Hs.2806 >10 R45596 Hs.3085 >10 R45566 Hs.3085 >10 R45596 Hs.39405 >10 R45367 Hs.3557 >10 R42031 Hs.1557 >10 R42031 Hs.3557	EST - RC_R09241		R09241	٧10	δ.
>10 R09241 >10 AA43738B Hs.115726 >10 U29953 Hs.715726 >10 AA39968B Hs.97663 >10 AA39968B Hs.97663 >10 U394B7 Hs.250 >10 AA400272 Hs.97758 >10 AA28228 Hs.118463 >10 AA28228 Hs.118463 >10 AA435901 Hs.56874 >10 AA435901 Hs.523841 >10 W93121 Hs.100924 >10 W73126 Hs.100924 >10 R74386 Hs.100924 >10 R74386 Hs.100924 >10 R74386 Hs.2037 >10 R74386 Hs.30905 >10 R74586 Hs.30850 >10 R74586 Hs.3037 >10 R74586 Hs.30357 >10 R74587 >10 R7477 Hs.89552		Hs.111301	AA482603	V10	æ
>10 AA482603 Hs.111301 >10 AA437368 Hs.111376 >10 U29953 Hs.76110 >10 AA339686 Hs.97669 >10 AA339686 Hs.97669 >10 AA339487 Hs.250 >10 U39487 Hs.27610 >10 AA400272 Hs.97758 >10 AA282238 Hs.127610 >10 AA28238 Hs.18633 >10 AA43801 Hs.26874 >10 AA4386 Hs.10624 >10 AA4386 Hs.10624 >10 AA45360 Hs.23841 >10 AA45360 Hs.2037 >10 AA45360 Hs.3080 >10 AA45580 Hs.3087 >10 AA45580 Hs.3087 >10 AA45580 Hs.3087 >10 AA45580 Hs.3085 >10 AA45580 Hs.3085		Hs.75092	R53966	×10	23
>10 R53966 Hs.75092 >10 AA422603 Hs.111301 >10 AA43784 Hs.115726 >10 AA437868 Hs. 187669 >10 AA439686 Hs.37609 >10 AA430968 Hs.37669 >10 AA430978 Hs.250 >10 AA43007 Hs.118463 >10 AA28223 Hs.118463 >10 NZ4879 Hs.36874 >10 NZ4879 Hs.10924 >10 AA435901 Hs.10924 >10 R74366 Hs.309405 >10 R73075 Hs.23650 >10 R74559 Hs.3550 >10 R745360 Hs.94550 >10 R745360 Hs.94550		Hs.107374	AA341723	210	o
>10 AA341723 Hs. 107374 >10 R53966 Hs. 11301 >10 R62366 Hs. 11301 >10 R09243 Hs. 11576 >10 AA437388 Hs. 11676 >10 U29953 Hs. 76110 >10 U29953 Hs. 7669 >10 U29953 Hs. 11576 >10 AA430968 Hs. 3766 >10 AA400212 Hs. 250 >10 AA400212 Hs. 118463 >10 AA282238 Hs. 118463 >10 AA282238 Hs. 118463 >10 AA282238 Hs. 118463 >10 AA438901 Hs. 58874 >10 AA438901 Hs. 10824 >10 R74366 Hs. 10824 >10 R74366 Hs. 10824 >10 R73075 Hs. 38046 >10 AA455960 Hs. 389405 >10 AA455960 Hs. 38955 >10 AA45380 Hs. 13857		Hs.74670	N57464	×10	4
>10 N57454 Hs.74670 >10 AA341723 Hs.107374 >10 AA482086 Hs.75092 >10 AA482083 Hs.11301 >10 AA437388 Hs.115726 >10 AA437388 Hs.76110 >10 U22953 Hs.76110 >10 AA439686 Hs.37758 >10 AA400212 Hs.250 >10 AA400212 Hs.17610 >10 AA400212 Hs.17660 >10 AA282238 Hs.118463 >10 AA282238 Hs.118463 >10 AA282380 Hs.10463 >10 WA31391 Hs.56814 >10 WA3136 Hs.106924 >10 WA3136 Hs.106924 >10 WA3136 Hs.106924 >10 AA455960 Hs.8068 >10 AA455960 Hs.309405 >10 AA455960 Hs.30405 >10 AA455960 Hs.30405 >10 </td <td></td> <td>Hs.17311</td> <td>AA487895</td> <td>×10</td> <td>75</td>		Hs.17311	AA487895	×10	75
>10 AA467695 Hs.17311 >10 N57464 Hs.17670 >10 R53966 Hs.175092 >10 AA487263 Hs.11301 >10 AA487388 Hs.115726 >10 AA4339686 Hs.76110 >10 AA4399686 Hs.97640 >10 U29953 Hs.18775 >10 AA4399686 Hs.97768 >10 AA4399686 Hs.97768 >10 AA439968 Hs.118463 >10 AA439968 Hs.118463 >10 AA439968 Hs.91756 >10 AA43991 Hs.250 >10 AA43991 Hs.13681 >10 AA43591 Hs.10634 >10 AA43591 Hs.10634 >10 AA43591 Hs.10634 >10 AA43591 Hs.106924 >10 AA45596 Hs.10692 >10 AA45596 Hs.10692 >10 AA45596 Hs.1069 >10	EST - RC_AA609907		AA609907	۷١٥	£
>10 AA669907 >10 AA487895 Hs.17311 >10 NS7464 Hs.17311 >10 AA341723 Hs.107374 >10 R53966 Hs.75092 >10 R6241 Hs.11301 >10 AA437388 Hs.111301 >10 AA437388 Hs.11526 >10 U28953 Hs.1610 >10 U28953 Hs.1756 >10 U39487 Hs.250 >10 AA39686 Hs.97669 >10 AA39686 Hs.11863 >10 AA39686 Hs.97669 >10 AA39686 Hs.11863 >10 AA400272 Hs.11863 >10 AA487591 Hs.58674 >10 R74879 Hs.10924 >10 R74369 Hs.10924 >10 R74369 Hs.10924 >10 R74369 Hs.10350 >10 R74369 Hs.10350 >10 R74369 Hs.10350<		Hs.98763	AA431797	, 10	9
>10 AA431797 Hs.98763 >10 AA487897 Hs.14670 >10 AA487893 Hs.14670 >10 NS7484 Hs.14670 >10 AA341723 Hs.107374 >10 R53966 Hs.75092 >10 R63963 Hs.111301 >10 AA43738 Hs.114726 >10 AA437386 Hs.16766 >10 AA439666 Hs.37669 >10 AA439666 Hs.18766 >10 AA43067 Hs.250 >10 AA40027 Hs.1863 >10 AA439966 Hs.11863 >10 AA43991 Hs.1861 >10 AA43591 Hs.1863 >10 R7486 Hs.10924 >10 R7486 Hs.10924 >10 R7486 Hs.10924 >10 R74590 Hs.3960 >10 R74596 Hs.3960 >10 R745960 Hs.9405 >10 R74596 </td <td></td> <td>Hs.99397</td> <td>AA455178</td> <td>>10</td> <td>œ</td>		Hs.99397	AA455178	>10	œ
>10 AA455178 Hs.99387 >10 AA411797 Hs.99387 >10 AA4879807 Hs.198763 >10 AA4879807 Hs.74670 >10 AA487486 Hs.74670 >10 AA482603 Hs.11301 >10 AA482603 Hs.11301 >10 RG9243 Hs.11301 >10 AA482603 Hs.11301 >10 AA437386 Hs.11576 >10 AA400212 Hs.266 >10 AA40021 Hs.267 >10 AA40021 Hs.11463 >10 AA40021 Hs.2681 >10 AA40021 Hs.2681 >10 AA40021 Hs.2681 >10 AA438901 Hs.118463 >10 AA438901 Hs.26814 >10 AA438901 Hs.2081 >10 AA438901 Hs.2081 >10 R7486 Hs.101 >10 R74866 Hs.101 >10 <th< td=""><td>_</td><td>Hs.99289</td><td>AA452606</td><td>×10</td><td>ន</td></th<>	_	Hs.99289	AA452606	×10	ន
>10 AA452606 Hs.99289 >10 AA4517B Hs.99397 >10 AA4609907 Hs.98763 >10 AA4609907 Hs.17311 >10 AA4809907 Hs.17311 >10 AA4809903 Hs.17314 >10 AA482606 Hs.17502 >10 AA482603 Hs.11301 >10 AA482606 Hs.11576 >10 AA482738 Hs.11576 >10 AA482968 Hs.1876 >10 AA492623 Hs.11465 >10 AA402238 Hs.11465 >10 AA402238 Hs.11465 >10 AA402601 Hs.56874 >10 AA435901 Hs.58874 >10 AA435901 Hs.10924 >10 Wa3121 Hs.2092 >10 Wa3121 Hs.2092 >10 Wa435901 Hs.30940 >10 AA455960 Hs.7557 >10 AA455960 Hs.39405 >10<		Hs.74566	078014	01 ^	_
>10 D78014 Hs.74566 >10 AA452605 Hs.99289 >10 AA45178 Hs.99397 >10 AA467895 Hs.17311 >10 AA487895 Hs.17311 >10 AA487895 Hs.17311 >10 N57464 Hs.7602 >10 R53966 Hs.7509 >10 AA437388 Hs.11301 >10 AA437388 Hs.11576 >10 AA4339686 Hs.11576 >10 AA4339686 Hs.11576 >10 AA4339686 Hs.11576 >10 AA4339686 Hs.11576 >10 AA439686 Hs.11643 >10 AA439686 Hs.11643 >10 AA439686 Hs.1043 >10 AA439691 Hs.26814 >10 AA43591 Hs.1043 >10 AA43591 Hs.106924 >10 AA43591 Hs.106924 >10 AA43591 Hs.106924 >10 <td>_</td> <td>Hs.84753</td> <td>D87433</td> <td>×10</td> <td>_</td>	_	Hs.84753	D87433	×10	_
>10 DB7433 Hs.84753 >10 DB7433 Hs.84753 >10 AA452606 Hs.99387 >10 AA451787 Hs.98783 >10 AA467897 Hs.174670 >10 AA487895 Hs.174670 >10 AA487895 Hs.174670 >10 AA487895 Hs.175034 >10 AA487895 Hs.11301 >10 AA487895 Hs.11301 >10 AA487895 Hs.11301 >10 AA487895 Hs.115726 >10 AA439686 Hs.11301 >10 AA439686 Hs.11301 >10 AA439686 Hs.11301 >10 AA439686 Hs.114663 >10 AA39686 Hs.118643 >10 AA39686 Hs.118643 >10 AA480531 Hs.250 >10 AA48291 Hs.36874 >10 AA48391 Hs.10924 >10 R74879 Hs.10924 >10	_	Hs.74669	W72859	×10	2
>10 W72859 Hs.74669 >10 DB7433 Hs.84569 >10 DB7433 Hs.84556 >10 AA45506 Hs.99289 >10 AA45178 Hs.99387 >10 AA487897 Hs.99389 >10 AA487897 Hs.17311 >10 AA487897 Hs.17311 >10 AA487897 Hs.17311 >10 AA487893 Hs.11301 >10 R6396 Hs.74670 >10 AA48789 Hs.11506 >10 AA48789 Hs.11576 >10 AA48066 Hs.16769 >10 AA48066 Hs.11576 >10 AA400212 Hs.11673 >10 AA40021 Hs.11643 >10 AA40021 Hs.356 >10 AA40021 Hs.3681 >10 AA435901 Hs.3681 >10 AA435901 Hs.10924 >10 R74879 Hs.10924 >10 R7	Basic fibroblast growth factor (bFGF) receptor (shorter form)	Hs.748	X66945	×10	60
>10 X66945 Hs.748 >10 W72859 Hs.7486 >10 D87433 Hs.84753 >10 D78014 Hs.7466 >10 AA452606 Hs.99387 >10 AA45178 Hs.99387 >10 AA45179 Hs.99387 >10 AA487985 Hs.17311 >10 AA487985 Hs.17311 >10 AA487986 Hs.74670 >10 AA48786 Hs.17301 >10 AA487603 Hs.11301 >10 AA48766 Hs.141301 >10 AA48766 Hs.141301 >10 AA48766 Hs.11406 >10 AA43738 Hs.114663 >10 AA43966 Hs.9693 >10 AA43966 Hs.9693 >10 AA43801 Hs.58874 >10 AA43801 Hs.18608 >10 AA43801 Hs.1010 >10 AA43801 Hs.10924 >10 AA4		Hs.50002	U77180	×10	'n
>10 UT7180 Hs.748 >10 X66945 Hs.748 >10 W72859 Hs.748 >10 W72859 Hs.7466 >10 D87431 Hs.7456 >10 AA452606 Hs.99289 >10 AA452780 Hs.99289 >10 AA45178 Hs.99289 >10 AA45178 Hs.99787 >10 AA487969 Hs.17311 >10 AA487969 Hs.17311 >10 AA487699 Hs.17311 >10 AA487699 Hs.11301 >10 AA487699 Hs.11301 >10 AA487699 Hs.11576 >10 AA48738 Hs.11576 >10 AA48738 Hs.11643 >10 AA400212 Hs.9663 >10 AA43996 Hs.118463 >10 AA438901 Hs.1863 >10 AA438901 Hs.9683 >10 AA438901 Hs.10924 >10 Wa4		Hs.10683	R45577	>10	œ
>10 R45577 Hs.10683 >10 VAR5845 Hs.10680 >10 VAR5854 Hs.74669 >10 WT2859 Hs.74569 >10 WAA52606 Hs.93397 >10 AA452760 Hs.93697 >10 AA45778 Hs.17311 >10 AA457897 Hs.17311 >10 AA487895 Hs.17311 >10 AA487895 Hs.17311 >10 AA487895 Hs.11301 >10 AA487895 Hs.11301 >10 AA487895 Hs.11301 >10 AA487895 Hs.11301 >10 AA487896 Hs.11576 >10 AA482738 Hs.11576 >10 AA433966 Hs.11663 >10 AA433966 Hs.1016 >10 AA438901 Hs.1663 >10 AA438901 Hs.10634 >10 AA438901 Hs.1064 >10 AA438901 Hs.1064 >10		Hs.20733	AA405199	>10	2
>10 AA405199 Hs. 2073 >10 RA4577 Hs. 10683 >10 X66945 Hs. 748 >10 X66945 Hs. 748 >10 X66945 Hs. 748 >10 W72859 Hs. 74653 >10 W72859 Hs. 74666 >10 AA452606 Hs. 99397 >10 AA452606 Hs. 99387 >10 AA451737 Hs. 10314 >10 AA487895 Hs. 174670 >10 AA487895 Hs. 174670 >10 AA487896 Hs. 174670 >10 AA487895 Hs. 11301 >10 AA43783 Hs. 11301 >10 AA43783 Hs. 11301 >10 AA43786 Hs. 114670 >10 AA43786 Hs. 11301 >10 AA43786 Hs. 11467 >10 AA43786 Hs. 11669 >10 AA43968 Hs. 11846 >10 AA43968 Hs. 1096 >		Hs.17998	AA243654	, 0	7

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fachykinin 2 (substance K neurokinin A neurokinin 2 neuromedin L neurokinin alpha neuropepiide K neuropepiide gamma)
                                                                                                                                                                                                             ESTs Moderately similar to protein phosphatase 1 binding protein PTG [M.musculus]
                                                                                                                                                                                                                                                                                                                      Small Inducible cytokine A5 (RANTES)
ESTS Highly similar to OSTEOINDUCTIVE FACTOR PRECURSOR (Bos taurus)
                                                                                                                                                                                                                                                                                                           ESTs Moderately similar to DNAJ PROTEIN HOMOLOG 1 [Homo sapiens]
                                                                                                                                                                                                                                   Homo saplans neural cell adhesion molecule (CALL) mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                                                                          Human protein tyrosine khase t-Rort (Ror1) mRNA complete ods
                      Homo sapiens mRNA for cardiac calsequestrin complete cds
                                                                                                              Homo sapiens clone 23798 and 23825 mRNA sequence
                                                                                                                                                                                                                                                                                                                                             PROSTATE-SPECIFIC MEMBRANE ANTIGEN
                                                                                                                         Human chemokine (TECK) mRNA complete cds
           SERUM AMYLOID A PROTEIN PRECURSOR
                                                                                                                                                                                                                                                                                                                                                                                                    Homo sapiens mRNA for smoothelin
                                                                                                                                                                                                                                                                             EST - HG3117-HT3293
                                                                                                                                                                                                                         ESTs
                                                                                                                                                                                                                                                                                        Hs.15903
Hs.62630
Hs.101393
                                                                                                                                                       Hs.13716
Hs.2563
Hs.65325
Hs.112629
Hs.55181
                    Hs.57975
Hs.3576
Hs.112087
                                                                                                                                                                                                                                                                                                                                                                                        Hs.26100
                                                                                                                                                                                                                                                                                                                         Hs.141503
                                                                                                                                                                                                                                                                                                                                   Hs.109439
                                                                                                                                                                                                                                                                                                                                                                             Hs.76487
Hs.17778
Hs.3157
                                                                                                                                                                                                                                                        Hs.125176
                                                                                                                                                                                                                                                                                                                                                         Hs.112065
                                                                                                                                                                                                                                                                                                                                                                                                     Hs.78483
                                                                                                                                                                                                                                                                                                                                                                                                                         Hs.1944
Hs.50652
                                                      Hs.42996
Hs.59486
                                                                                                                                                                                                              Hs.12112
                                                                                                                                                                                                                         Hs.47438
                                                                                                                                                                                                                                   Hs.21226
                                                                                                                                                                                                                                              Hs.43148
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                                                                                                                                    Hs.43125
                                                                                                   Hs.29653
                                                                                                              Hs.6326
                                                                             Hs.60162
                                                                                        Hs.33455
                                                                                                                        Hs.50404
                                                                                                                                             Hs.38022
                                                                                                                                 AA443800
AA136353
AA284920
AA446659
T16335
AA69018
N95796
H89980
N5254
AF002246
N75215
AA486185
H59887
                                                                                                                                                                                                                                                                           HG3117-
T85315
AA043349
AA434108
F02702
D62584
                                                                                                                                                                                                                                                                                                                                            M99487
AA435805
U66061
R06986
R54179
                     AA055163
AA234383
AA401404
AA045306
                                                                            AA005236
R85880
R77493
T16211
                                                                                                                                                                                                                                                                                                                                                                                                    Y13492
                                                                                                                        U86358
                                                                  W93497
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153		TS:	EST	ESTs	ES33	ESTs	ES13	ESTS	EST	Human mRNA for KIAA0278 gene partial cds	ESTs	ESTS	EST	ntegrin beta 2 (antigen CD18 (p95) tymphocyte function-associated antigen 1 macrophage antigen 1 (mac-1) heta subunit)	Homo sapiens telomeric repeat binding factor (TRF1) mRNA complete cds	ESTs	ESTS	ENTS	Homo sapiens ribonuclease P protein subunit p20 (RPP20) mRNA complete cds	Transient receptor potential channel 1	ESTS	Human YMP mRNA complete cds	ESTs	ESTS	ESTs Weakly similar to unknown protein [H.saplens]	ESTs	EST - RC_N63688	Solute carrier family 3 (cystine dibasic and neutral amino acid transporters activator of cystine dibasic and neutral amino acid transport	ESTS	ESTs	ESTs Weakly similar to centaurin alpha (R.norvegicus)	ESTs Moderately similar to PANCREATITIS-ASSOCIATED PROTEIN 1 PRECURSOR [H.sapiens]	Human NECDIN related protein mRNA complete cds	BRAIN SPECIFIC POLYPEPTIDE PEP-19	ESTs	Thrombopoletin (myetoproliferative leukemia virus oncogene ligand megakaryocyte growth and development factor)	Homo sapiens short form transcription factor C-MAF (c-mat) mRNA complete cds	ESTs	ESTs
25,478	Hs 71719	4s.112737	Hs.60418	ls.122656	Hs.86045	Hs.15342	Hs.7120	Hs.8769	4s.139171	Hs.40888	Hs.23213	Hs.14898	4s.104249	Hs.83968	Hs.90357	Hs.5476	Hs.32246	Hs.116428	Hs.18747	Hs.94413	Hs.98998	Hs.9999	Hs.28137	Hs.66357	Hs.18767	Hs.124952		Hs.110	Hs.6952	Hs.86693	Hs.28802	Hs.112961	Hs.50130	Hs.80296	Hs.87469	Hs. 1166	Hs.30250	Hs.144599	Hs. 109896
0002500	AA142875	-	AA010611	W15376 P	AA196306	W15580	T23457	AA428258	AA227469	D87468	H17865	N92824	AA233380	X64072	U74382	T32561	AA235009	W31906	AA203527	X89066	AA443311	U52101	H27852	C14270	H57357	Z39300	N63688	H92451	T17215	AA480886	AA262556	AA399061	U35139	U52969	AA235984	L36051	AA043501	AA291983	W44682
,	5 5	5 2	>10	5,	5	210	5,	5	5	×10	, 10	V 10	5	×10	210	٧10	٠ ٢	710	v 10	۲٠ د ۲۵	۲٠ د ۲۰	, to	, 10	×10	ot <	5	5	, 10	\$, 10 10	>10	ot	ě	× 10	×10	×10	۰10 م	5	v 10
90700	10406	38939	16206	32810	18210	24054	23047	12944	34172	9317	19331	21035	34208	5974	5032	41941	34239	32852	7662	6432	37001	4630	19489	28483	19801	24672	31153	40250	23028	28072	11868	35359	4285	4655	26030	2042	25262	34821	42405

41348	710	R40395	Hs.112125	Lecithin-cholesterol acyltransferase
14494	>10	AA609645	Hs.25632	ESTs Weakly similar to PNG gene [H.sapiens]
4453	, 10	U43916	Hs.79368	Human epithelial membrane protein (CL-20) mRNA complete cds
42758	×10	Z41411	Hs.107040	ESTs
2098	>10	L39009		EST - L39009
35637	, 6	AÁ402933	Hs.29283	ESTs
16549	×10	AA029697	Hs.94854	ESTs
1220	×10	HG3733-		EST - HG3733-HT4003
39934	×10	H52185	Hs.124994	ESTs
7735	×10	AA232121	Hs.109631	Human tyrosyl-IRNA synthetase mRNA complete cds
40392	6	H99587	Hs.108880	ESTs
37170	>10	AA447779	Hs.99145	EST
18361	^ 10	AA223902	Hs.86899	ESTs
19366	۷. ۲0	H19204	Hs.133466	ESTs
38429	>10	AA496965	Hs.108694	Glycophorin A
38021	>10	AA481059	Hs.105152	ESTs
5184	> 10	U82169	Hs.87234	Human frizzled homolog (FZD3) mRNA complete cds
27863	×10	AA458923	Hs.70202	ESTs Weakly similar to F23B2.4 (C.elegans)
37476	٧١٥	AA455051	Hs.99386	EST
859	٧10	D87468	Hs.40888	Human mRNA for KIAA0278 gene partial ods
27185	۷10	AA410895	Hs.62348	ESTs
41010	v 10	N80686	Hs.21639	Human APEG-1 mRNA complete cds
38241	40	AA489076	Hs.105101	ESTs
22701	۷10 م	R89477	Hs.34299	ESTs
12152	×10	AA291271	Hs.10886	ESTs Weakly similar to uroporphyrinogen III synthase UROIIIS [H.sapiens]
38913	۷10	AA609531	Hs.112050	ESTs
34034	۷10	AA192871	Hs.83760	Troponin I (skeletal fast)
37644	۰10	AA459857	Hs.99503	EST
4173	01.	U29725	Hs.3080	Human BMK1 alpha kinase mRNA complete cds
16178	>10	AA009839	Hs.1632	CD27L RECEPTOR PRECURSOR
20527	>10	N54161	Hs.124044	ESTs
41918	٠10 ۲	T25873	Hs.102243	ESTs
1525	×10	J04621	Hs.1501	Syndecan 2 (heparan sulfate proteoglycan 1 cell surface-associated fibroglycan)
19160	٧١٥	H10208	Hs.30972	EST
16860	×10	AA055833	Hs.58152	ESTs Weakly similar to Natsu [M.musculus]
36927	>10	AA437259	Hs.104944	EST
1595	01.4	K03207	Hs.103972	Salivary proline-rich protein
15574	, 10	W38778	Hs.26216	ESTs
8985	10	C00125	Hs.24332	ESTs Weakly simitar to similar to deoxynbose-phosphate aldolase [C.elegans]
33995	۰10 م	AA182845	Hs.139088	Homo sapiens FIP2 afternatively translated mRNA complete cds

ESTS	EST\$	Homo sapiens PAC done DJ130H16 from 22q12.1-qter	ESTs	ESTRADIOL 17 BETA-DEHYDROGENASE 1	duman protein tyrosine phosphatase PTPCAAX1 (hPTPCAAX1) mRNA complete cds	ESTs	ESTS	EST - RC_T59537	Apolipoprotein C-III	-tydroxyacyt-Coenzyme A dehydrogenase/3-ketoacyt-Coenzyme A thiolase/enoyt-Coenzyme A hydratase (trifunctional protein)	H.saplens mRNA for fibrinogen-like protein (pT49 protein)	Homo sapiens mRNA for KIAA0673 protein partial cds	Lymphotoxin-beta	Homo saplens mRNA for KIAA0679 protein partial cds	ESTs	EST - RC_AA48334	ESTs	ESTs .	ESTs	ESTs	ESTs	ESTs	Prostagiandin E receptor 3 (subtype EP3) (alternative products)	ESTs Weakly similar to !!!! ALU SUBFAMILY SX WARNING ENTRY !!!! [H.sapiens]	ESTs	ESTS	ESTS	Homo sapiens DBI-related profein mRNA complete cds	EST	ESTS	ESTs	ESTs	Homo sapiens mRNA for GABA-BR1a (hGB1a) receptor	ESTs	ESTs Highly similar to FIBROPELLIN C PRECURSOR (Strongylocentrotus purpuratus)	ESTs	EST - RC_F12567	CD20 RECEPTOR	ESTs Weakly similar to GLYCEROPHOSPHORYL DIESTER PHOSPHODIESTERASE (E.coll)
Hs.34956		Hs.25199 +	Hs.7858 E	Hs.85279 E	_	Hs.25209 E	Hs.96744	_	Hs.73849 /	Hs.75860	Hs.2659	Hs. 106487	Hs.890	Hs.5734	Hs.97514	_	Hs.57929	Hs.50891	Hs.12701	Hs.22505	Hs.46987	Hs.97899	Hs.495	Hs.29759	Hs.61307	Hs.49051	Hs.124953	Hs.15250	Hs.97250	Hs.6598	Hs.47566	Hs.124964	Hs.14829	Hs.7974	Hs.21041	Hs.95511		Hs.89751	Hs.107755
AA283620	AA034918	AA252191	AA092348	U34879	T40895	R54594	AA419011	T59537	172491	D16480	Z36531	R42233	AA287870	W01875	AA421158	AA448334	AA448625	N80279	238289	F03111	AA037433	AA469952	X83857	C01833	AA025728	AA400102	240646	AA399269	AA291522	H89355	N53444	R81949	AA348198	AA495865	AA453034	AA085721	F12567	X07203	R82942
v.	×10	٧10	55	>10	×10	۲٠ د ۲۵	6,	01.¢	5	>10	۲۰ د د	×10	×10	×10	٧10	>10	>10	510	>10	×10	5	01 <	٧١٥	>10	۰ to	01 %	×10	۰10 م	×10	01^	>10	01 ^	×10	>10	, 0	10	v.	×10	۸ ک
7949	16607	11670	7354	4277	23214	22209	36151	23372	42136	289	15974	41379	34764	24027	36197	37211	27684	31790	24515	18652	16635	37815	6364	9034	16469	27034	42746	35368	34805	19983	31126	22616	12246	1118	13486	25512	29073	5541	41689

PESTS Hemoglobin gamma-G EST ESTS ESTS SESTS Highly similar to THREONYL-TRNA SYNTHETASE CYTOPLASMIC [Homo sapiens] ESTS ESTS Highly similar to FORMYLTETRAHYDROFOLATE DEHYDROGENASE [Raitus norvegicus] Pyruvate carboxylase ESTS COTP synthetase ESTS COTP synthetase ESTS COTP SYNTHETASE ESTS COTP SYNTHETASE ESTS ESTS COTP SYNTHETASE ESTS ESTS ESTS			Homo sapiens Ca2+-dependent phospholipase A2 mRNA complete cds 1 ESTs Glycoprotein Ib (platelet) beta polypeptide 1 CYCLIN-DEPENDENT KINASE INHIBITOR 1 2 Lymphocyte cytosolic protein 1 (L-plastin) 3 Lymphocyte cytosolic protein 1 (L-plastin) 4 AFFX-TrpnX-5 1 Macrophage stimulating 1 (hepatocyte growth factor-like) 5 ESTs
Hs. 125052 Hs. 112155 Hs. 89554 Hs. 88602 Hs. 12382 Hs. 14238 Hs. 14238 Hs. 107365 Hs. 107365 Hs. 89890 Hs. 89890 Hs. 29924 Hs. 2693 Hs. 26953 Hs. 26553 Hs. 26553	Hs. 78362 Hs. 27261 Hs. 31604 Hs. 6923 Hs. 104423 Hs. 30343 Hs. 30343	Hs. 16611 Hs. 76224 Hs. 98450 Hs. 24587 Hs. 144323 Hs. 144323 Hs. 46146	Hs.290 Hs.30484 Hs.3847 Hs.74984 Hs.76506 Hs.76034
R80965 AA490916 R92458 AA434246 C14784 R86970 R52163 AA086487 N3867 H17611 AA404494 D59722 X82694 AA466694	N64344 H09343 H18706 AA121534 AA243574 AA521370 R22139 N26740	U44429 U03877 AA426056 N50550 K02100 T64891 AA418001	U03090 H04768 U59632 U09579 J02923 AFFX- U37055
	6 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0		5 5 5 5 5 5 5 5
32343 38335 41729 36707 26491 41702 32246 17314 30325 19823 42153 19321 27110 27110 27110 27110 3333 37679	. 40829 19132 19353 10935 11621 38538 10095 30014	4464 3650 36377 20437 1576 42076 27257 30582	3631 19026 4752 3768 1437 33805 4310

TO SEE SEE SEE SEE SEE SEE SEE SEE SEE SE	45. 119218 45. 119218 45. 119348 45. 119348 45. 11936 45. 11936 45. 11936 45. 11936 45. 11936 46. 11936 47. 11936	X06256 AA457409 AA457409 AA46080922 AA6080988 D16532 AA713168 AA044732 AA046731 AA040379 C21481 U07620 X77307 T80833 N78765 U8979 R94521 H12674 H61046 AA449521	Representation of the property	37571 338800 27952 27621 38784 291 18014 16720 25336 2547 39953 13777 13419 1403 42373 21520 9796 33650 3772 42473 25195 25195 3772 6214 23575 31775 31775 31776 42473 25195 3712 6214 23575 31776 6214 23575 31776 3177
 ESTs PROTEIN IN PRP16-SRP40 INTERGENIC REGION [S.cerovisiae] 	Hs.26320	R43980	01.4	21911
	Hs.8108	T34622	×10	22184
		-	2	23184
•	_	AA011305	×10	1600
	•	3	2	16225
29 Cathepsin C	Hs.10029	AA011305	×10	30007
	Hs.8108	T34622	×10	
	113.40260	K43300	5	1911
	00030		:	3/200
	Hs.98428	AA449424	>10	37756
	-	AA489210	5,	38248
	1	90000	: 1	23410
	Hs.36702	H77734	01.4	9000
	HS.4044	T33511	>10	41960
	7707		?	1001
	Hs.4811	AA147537	410	10011
-	Hs.70405	H61046	۲ د	93266
_	MS.9390	H12674	۰10 د	9377
_			2	69/7
	Hs.124693	R94521	65.	
	HS.67840	082979	×10	5206
			2	1775
	Hs.50847	N79765	,	
	HS.14/34	T80833	710	3575
	10.4.01	2027	or <	3214
	He 2507	77207	•	4
	Hs.89661	U07620	×10	713
	HS.84630	C21481	, 5	8607
	2000	AA033730	210	5195
	He 75736	00755044		2
	Hs.122531	W63731	×10	F.473
	HS.106900	Z41239	0×	1812
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	11. 403466		2	200
	Hs.22631	AA404271	4	3 8
	Hs.73974	M63509	٠10	96/
	13.646.00	KZ0Z0/	220	220
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	Hs.103012	W35362	۲. د	373
	18.9555	300123	70	ន
DROPENKEPHALIN A PRECURSOR	U. 03557		2	P.
EST 3	Hs.22269	AA450336	×10	9
	2000.50	AA463304	×10	11
	CA05.01		2 :	3
	Hs.108144	H56010	×10	č,
	25.100	609CZW	×10	47
	4000	00000		2
	Hs.101404	AA053405	۰ 10	36
-	HS.//200	AA044/32	۲٠ د ۲٥	8
101	00022	200	2	4
ESTs Weakly similar to weakly similar to meakly similar to meave or the state of th	Hs.57672	AA173168	9,7	•
Very fow defining income and a second for the secon	HS./3/29	D16532	٠ و	_
in the second of	13.500	AAGUGAGG	×10	¥
Testis specific protein Y-linked	He 2051	80000044	•	;
FSIS	Hs.56589	AA446242	^10	7
	11S.47.042	AA463700	× 05×	22
Lamb sations CD39L3 (CD39L3) mRNA complete cds	11. 170.43	002001	2	2
EST	Hs.112636	A A 509052	,	. ,
TST.	Hs.99458	AA457409	710	
		X06256	, 10	_

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Homo saplens killer celt receptor (KIR103) mRNA allete ASDJ complete cds
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                                                                                                                                                                                EST • HG2416-HT2512
ESTs Moderately simllar to alfa subunit [H.sapiens]
                                                                                                                                   Complement component 8 gamma polypeptide
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               EST - RC_AA621750
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ESTs
EST - RC_R98947
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AA404219
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N71371
AA074407
N49308
AA4255485
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X15357
AA266946
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Tyrosinase (oculocutaneous albinism IA) ESTS Highly similar to ADENYLOSUCCINATE SYNTHETASE MUSCLE ISOZYME [Mus musculus]
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ESTs Weakly similar to precursor of major fibrous sheath protein [M.musculus]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human transductn-like enhancer protein (TLE3) mRNA complete cds
                                                             Homo saplens mRNA for zinc finger protein FPM315 complete cds
BASIC TRANSCRIPTION FACTOR 62 KD SUBUNIT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            S100 calclum-blnding protein A5 (formerly S100D)
                                                                                                                                                                                                                                                                                                                                                                                                                                      Human mRNA for KIAA0306 gene partial cds
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Glucocorticoid receptor	Carbamoyl-phosphate synthetase 1 mitochondrial	ESTs	TRANSFORMING PROTEIN RHOB	ESTs	EST - RC_AA255523	ESTs	Glycerol kinase 2 (testis specific)	Mannose-6-phosphate receptor (cation dependent)	EST - S78774	ESTs	ESTs	ESTs	Allograft Inflammatory factor 1	ESTs	EST	ESTs Weakly similar to RTP60 [R.norvegicus]	EST	ESTs	ESTs	ESTs	ESTs	EST	EST	Pleiotrophin (heparin binding growth factor 8 neurite growth-promoting factor	ESTs	ESTs	ESTs	ESTs Highly similar to 65 KD YES-ASSOCIATED PROTEIN [Gallus gallus]	ESTs	ESTS	ESTs	ESTs	ESTs	Hippocaldn-like 1	ESTs		ESTs	_	ESTs
Hs.75772	Hs.50966	Hs.17749	Hs.75122	Hs.42658		Hs.61555	Hs.98008	Hs.75709		Hs.25717	Hs.58550	Hs.20945	Hs.76364	Hs.98189	Hs.99489	Hs.126270	Hs.112591	Hs.112238	Hs.12610	Hs.6202	Hs.97450	Hs.89267	Hs.59332	Hs.44	Hs.110128	Hs.87762	Hs.55062	Hs.71873	Hs.22906	Hs.7915	Hs.111223	Hs.69009	Hs.98378	Hs.3618	Hs.20887	Hs.104425	Hs.92350	Hs.4188	Hs.36030
AA234527	T59148	T96123	AA452158	AA463434	AA25553	AA029428	X78712	AA393666	S78774	N68830	W79698	H06371	U19713	AA417063	AA459662	R62313	AA608792	AA470135	R11157	N22006	AA400795	AA284067	W90735	M57399	W88426	AA250845	N94581	AA148213	R44949	N79674	N51105	AA131919	AA423970	D16227	AA243598	AA279391	H01428	W46947	H65942
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34231	42046	23913	37333	27946	34407	16542	6248	8227	3507	40907	33340	19079	3992	36059	37634	41581	38734	. 37836	21303	20125	35516	26771	33558	2830	42625	26152	31988	17763	21959	10000	30658	17629	36260	285	26123	34535	29100	24122	19894

	Y PROTEIN RCA1 [Saccharomyces cerevisiae]		
H. sapiens CHML mRNA H. sapiens CHML mRNA Human preprocarboxypeptidase A2 (proCPA2) mRNA complete cds Epidermal growth factor receptor ESTs EST H. sapiens mRNA for cylich II ESTs ESTs	ESTs ESTS Highty similar to MITOCHONDRIAL RESPIRATORY CHAIN COMPLEXES ASSEMBLY PROTEIN RCA1 [Saccharomyces cerevisiae] EST - HG2260-HT2349 ESTs ESTs ESTs EST - HG2705-HT2801 EST - RC_T91283 Human APEG-1 mRNA complete cds ESTs	ESTs ESTs ESTs ESTs ESTs ESTs ESTs ESTs	ES18 Homo sapiens mRNA for KIAA0525 protein partial cds ESTs Homo sapiens putative transmembrane protein (CLN5) mRNA complete cds Treacher Collins syndrome susceptibility protein ESTs ESTs ESTs
HS.25420 HS.34514 HS.89717 HS.7432 HS.14627 HS.58663 HS.3232 HS.2332 HS.9567 HS.9657	Hs. 72639 Hs. 29385 Hs. 105618 Hs. 137530 Hs. 21639 Hs. 144212	Hs. 24872 Hs. 142462 Hs. 17404 Hs. 105686 Hs. 88888 Hs. 29279 Hs. 12420 Hs. 54643	HS.30309 HS.30379 HS.30213 HS.73166 HS.46784
T16258 X64728 W60008 X00588 AA034366 W81607 Z46788 H117618 R41836	AA166917 AA598437 HG2260- T79638 AA488997 HG2705- T91283 U57099 N66796	AA399633 AA279662 N39584 AA236868 AA488659 AA235874 U71207 U66162 S78467 AA506136 N90668 R01398 U89995 S81957	AA443956 AA036753 AA010328 N24772 W93015 AA418392 AA402000
			5 5 5 5 5 5
22987 5985 42461 5422 16603 33389 6931 19324 32166	17958 38569 1006 23552 38228 1066 23815 4699 31306	12389 34539 20358 26070 38210 26025 4978 4935 3501 14281 31859 41104 5293	27615 7152 16197 20176 33586 27265 12453

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ESTs Moderately similar to PUTATIVE SERINE/THREONINE-PROTEIN KINASE PKWA [Thermomonospora curvata]
                                                                                                                                                                     Human adenyty cyclass-associated protein homolog CAP2 (CAP2) mRNA complete cds Homo saplens mRNA for KIAA0515 protein partial cds
                                                                                                                                                                                                                                                                                                    Homo sapiens Pig12 (PIG12) mRNA complete cds
                                                                                                                          Chromogranin A (parathyroid secretory protein 1)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ESTs Moderately similar to FOG [M.musculus]
ESTs
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Homo sapiens done 23579 mRNA sequence
ESTs Weakly similar to WWP2 [H.sapiens]
                                                                                                                                                                                                                                                                                                                                                                                                                                 EST - M27533
                                                                                                                                                                                                                                                                                                                                                                                  ESTs
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Hs.106309
Hs.18397
                                                                                                                                                                                                                                                                                                                                                                                                                                                       Hs.8261 · Hs.83466 Hs.103102 Hs.46584
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Hs.30390
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Hs.119190
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Hs.72447
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H94043
H46167
H46167
M270305
F10265
M27533
AA082171
H28566
N33586
N64191
N23009
R54416
AA282583
T16473
AA48004
AA168173
R4449
AA168173
R4449
AA168173
F2625317
T96259
T15829
W27301
H03299
F10338
AA435896
  A4424652
T40448
A4114071
T67026
Z41697
T56470
A4424331
A4424331
A4424331
H97638
A4160530
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Human pre-TANK cell associated protein (1D12A2) mRNA complete cds		Laminin gamma 1 (formerly LAMB2)	Refinoblastoma-binding protein 1(alternative products)	: ESTs	Myasthenic syndrome antigen B (human fetal brain mRNA 3477 nt)	ESTs	ESTs	ESTs Weakly similar to keratin 8 type II cytoskeletal embryonic (M.musculus)	_	CELL DIVISION PROTEIN KINASE 8	ESTs	ESTs	ESTs	Homo sapiens transmembrane protein mRNA complete cds	EST	EST - U38372	Human MHC Class I region proline rich protein mRNA complete cds	3 Human Meis1-related protein 2 (MRG2) mRNA partial cds	ESTs	EST) EST	EST - M11591	ESTs	3 EST	Human clones 23920 and 23921 mRNA sequence	Human U1-snRNP binding protein homolog mRNA complete cds	ESTs		_		EST - RC_R92512_s		EST	ESTs	Human two P-domain K+ channel TWIK-1 mRNA complete cds	ESTs	EST - RC_AA101056	EST - AF001359_f	Human msg1-related gene 1 (mrg1) mRNA complete cds
Hs.278	Hs.25536	Hs.87428	Hs.91797	Hs.101248	Hs.30941	Hs.75169	Hs.87298	Hs.16003	Hs.128630	Hs.25283	Hs.65973	Hs.50429	Hs.96837	Hs.110903	Hs.33416		Hs.41548	Hs.117313	Ms.47606	Hs.48382	Hs.104059		Hs.23017	Hs.112603	Hs.7571	Hs.93502	Hs.99043	Hs. 105042	Hs.22646	Hs.27278		Hs.47390	Hs.59890	Hs.9410	Hs.79351	Hs.93675			Hs.82071
L17325	C17938	M55210	\$57153	T26444	U95019	R44234	AA262972	W24127	AA232251	X85753	Z40689	W87484	AA344854	AA621414	R83664	U38372	AA456966	U68385	N53043	N59432	AA181935	M11591	W84413	AA608852	U79271	U44798	AA446000	AA459392	R44477	F03889	R92512	N51987	AA001879	T52201	030065	AA148923	AA101056	AF001359	U65093
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1850	9101	2807	3383	23142	5367	32205	26515	15332	34193	6392	33784	33474	34964	14584	22640	4339	37557	4937	30795	30966	33991	2265	24315	38752	5119	15037	37045	37627	21935	18669	22737	30727	16086	23293	5294	17769	25549	\$	4856

9.1/236

ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	EST - RC_T94409	EST	ESTs	EST - RC_H61560	EST - HG831-HT831	Human mRNA for rod photoreceptor protein complete cds	Human metabotropic glutamate receptor 8 mRNA complete cds	GUANYLATE CYCLASE SOLUBLE BETA-1 CHAIN	ESTs	ESTs	EST - RC_AA448226	ESTs	Diacylglycerol kinase alpha (80kD)	ESTs	ESTs	ESTs	EST	ESTs	ESTs	ESTs Weakly similar to putative type III atcohol dehydrogenase (O.melanogaster)	Gonadotropin-releasing hormone (teutinizing-releasing hormone)	EST	ESTs	ESTs	Homo sapiens G protein beta 5 subunit mRNA complete cds	ESTs	STERYL-SULFATASE PRECURSOR	EST .	ESTs Weakly similar to CLATHRIN COAT ASSEMBLY PROTEIN AP17 [Rattus norvegicus]	ESTs	Homo sapiens Grb14 mRNA complete cds	ESTs	ESTs Weakly similar to putative progesterone binding protein [H.sapiens]	GRANZYME A PRECURSOR
Hs.107256	Hs.98416	Hs.57489	Hs.98983	Hs.16446	Hs.20526		Hs.58009	Hs.21107			Hs.26886	Hs.86204	Hs.77890	Hs.26026	Hs.26615		Hs.108509	Hs.74044	Hs.60992	Hs.41585	Hs.20423	Hs.71647	Hs.13121	Hs.124694	Hs.11553	Hs.82963	Hs.96869	Hs.14480	Hs.8861	Hs.115241	Hs.99598	Hs.79876	Hs.49112	Hs.50628	Hs.59163	Hs.83070	Hs.23786	Hs.109494	Hs.90708
AA214730	AA424535	AA125781	AA442779	AA010619	R11654	T94409	W69435	AA479299	H61560	HG831-	D63813	U92459	X66533	R53972	Z38900	AA448226	AA250836	X62535	AA019603	AA609080	H98854	AA136541	H98768	N74604	R63545	H87229	AA347417	179203	AA421778	AA017518	AA463627	M16505	N56062	N75507	AA598959	L76687	AA400292	N92882	M18737
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7697	36296	17490	36976	10425	41196	42285	33185	13974	19868	1378	621	5346	6209	22196	24609	37196	26151	5938	16373	28356	20070	17717	20069	40985	10194	40200	34983	23543	12770	25085	37746	2370	31244	31716	28306	2220	12404	41050	2407

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Small Inducible cytokine A5 (RANTES)
Spleen focus forming virus (SFFV) proviral integration oncogene spi1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           H.sapiens mRNA for F25B3.3 kinase like protein from C.elegans
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Homo sapiens transmembrane protein mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      EST Weakly similar to precursor polypeptide [H.saplens]
                                                                                                                                                                     ESTs Weakly similar to P24 protein [M.musculus]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         PLATELET GLYCOPROTEIN V PRECURSOR
                                                       Homo sapiens CAG-isf 7 mRNA complete cds
                                                                                                                Suppression of tumorigenicity 2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             CYTOCHROME P450 IVF3
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EST
EST
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Hs. 12992
Hs. 107614
Hs. 104010
Hs. 104010
Hs. 102923
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Hs. 23084
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Hs. 147503
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Hs.99235
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Hs.103668
Hs.55778
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T66948
N32060
AA445949
O12763
N93008
AA084405
R42039
AA233151
AA233151
AA42812
H90133
AA62041
AA260413
AA62041
AA260413
AA260241
AA260241
AA260241
AA260241
AA260241
AA260241
AA260241
AA260241
AA26024
R15890
R79239
AA461106
AA4610639
  23462
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41064
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Homo saplens glycogenin-2 gamma (glycogenin-2) mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human interleukin-11 receptor alpha chain mRNA complete cds
                                                                       Human butyrophilin (BTF1) mRNA complete cds
                                                                                                                                                                         Human mRNA for KIAA0369 gene complete cds
Zinc finger protein 135 (clone pHZ-17)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ESTs Weakly similar to ZK792.1 (C.elegans)
                                                                                                                          Homo sapiens nkat7 mRNA complete cds
                                                                                                                                                                                                                                                                        EST - RC_AA079094
                                                                                                                                                                                                                                                                                                                                                EST - RC_AA069386
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Hs. 95898
Hs. 12354
Hs. 48614
Hs. 58585
Hs. 10476
Hs. 106879
Hs. 26159
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Hs. 62781
Hs. 109610
Hs. 21355
Hs. 21358
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Hs. 58187
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Hs.87306
Hs.99539
Hs.54421
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Hs.57760
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Hs.55015
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Hs.76852
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AA233277
W57862
AA157772
U90543
AU90543
AU5081
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AA078094
W88568
AA430539
AA078098
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AA078098
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AA400888
AA401830
U32324
AA465650
AA401253
AA401253
AA401253
AA401253
AA431868
F10640
N62724
W79524
T59005
AA37841
R41389
N63965
S72370
AH17463
  11427

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330858

8870

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Dihydrolipoamide dehydrogenase (E3 component of pyruvate dehydrogenase complex 2-oxo-glutarate complex branched chain keto acid dehydrog
                                                                                                                                                                                                                                                                                                     ESTs Moderately similar to NITRIC OXIDE SYNTHASE INDUCIBLE [H.sapiens]
                                                                                                    Homo sapiens zinc-finger helicase (hZFH) mRNA complete cds
ESTs Highly similar to transmembrane receptor [M.musculus]
ESTs
                                                                                                                                                                                                                 Homo sapiens NF-E2 protein (NF-E2) mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                 Homo sapiens inRNA for KIAA0688 protein complete cds
                                                                                                                                                                                                                                                                                                                                                                            ESTs Highly similar to mosaic protein LR11 [H.sapiens]
                                                                                                                                                   Homo sapiens clone 24818 mRNA sequence
ESTs
EST - RC_AA079331
                                                                                                                                                                                                                                                                                                                                                                                                    EST - RC_AA457023
EST
ESTs
                                                                                                                                                                                                                                                                EST - RC_T82307
EST - RC_T98262
                                                                                       EST - RC_H03358
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Hs. 47566
Hs. 66180
Hs. 104005
Hs. 30494
Hs. 93677
                                                Hs.74635 E
Hs.19235 E
Hs.100530 E
                                                                                                                                                                                       Hs.34492
Hs.100472
Hs.75643
Hs.112890
Hs.43590
Hs.34882
                                                                                                                                                                                                                                                                                                                 Hs.89310
Hs.50446
Hs.25870
                                                                                                              Hs. 62633
Hs. 102329
Hs. 35598
Hs. 106823
Hs. 108107
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Hs.93996
Hs.60887
                                                                                                                                                                                                                                                                                          Hs.95111
Hs.97357
 Hs.24032
Hs.22971
Hs.76798
Hs.39252
                                                                                                   Hs.25601
                                                                                                                                                                                                                                                                                                                                                     Hs.42829
R26141
A4101632
N8666
N52398
A453435
A445475
A445472
W87423
C01394
N5268
AA073331
AN0305
AA073331
AN0305
AA468273
S77763
AA468273
AA468272
AA4840
D60364
T82307
T98262
AA453472
AA484181
AA284181
AA487023
T10134
N50666
H98700
A4457023
T50662
N62200
A445702
                                                                                       21481
10887
31431
30756
35829
24540
37500
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ESTs Weakly similar to dual specificity phosphatase [H.saplens]
ESTs Highly similar to S-ACY1. FATTY ACID SYNTHASE THIOESTERASE MEDIUM CHAIN (Rattus norvegicus)
                                                                                                                                                                                                                                                                                          Homo sapiens Shab-related delayed-rectifier K+ channel alpha subunit (KCNS3) mRNA complete cds
                                                                                                                                                                                                                                                                                                                                    ESTs Highly similar to RAS-RELATED PROTEIN RAB-14 (Rattus norvegicus)
                                                                                                                                                                                                                                                         ESTs Weakly similar to isopentenyl-diphosphate Delta-isomerase [H.sapiens]
                                                                                                                                                                                                                                                                               ESTs Weakly similar to !!!! ALU CLASS B WARNING ENTRY !!!! [H.sapiens]
           Human repressor transcriptional factor (ZNF85) mRNA complete cds
                                                                                                                                                                                                                                                                     ESTs Highly similar to PROTEIN CDC27HS [Homo saplens]
                                                                                                                                                                          Homo sapiens BAC clone RG113D17 from 7p14-p15
                                                                                            ESTs Weakly similar to unknown [S.cerevisiae]
                                                                                                                                                                                                                                                                                                                                                            H.sapiens TTF mRNA for small G protein
Homo saplens mRNA for Hic-5 partial cds
                        Human metallothlonein (MT)I-F gene
                                                                                                                            EST - RC_N49259
EST - HG2139-HT2208_f
                                                                                                                                                                                                                                                                                                                                                 Bradykinin receptor B2
                                                                                EST - RC_AA069920
                                               EST - RC_AA039568
                                                                                                                                                                                                                                                                                                                            EST - RC_T87648
                                                                                                      EST - RC_N34457
                                                                                                                                                              EST - RC_N21461
                                                                                                                                                                                                                                                                                                                                                                                                                                EST - D38462
                                                          ESTs
                                                                                                                                                    EST
                                                                                                                                                                                                                                                                                                                                     Hs.125175
Hs.54421
Hs.109918
Hs.25511
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Hs.63559
Hs.36372
Hs.47584
Hs.109870
Hs.18357
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Hs.24309
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 Hs.88756
Hs.37138
                        Hs.110440
                                                                                          Hs. 10839
                                                                                                                                                                                                                                              Hs.11722
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                                                                     Hs.44600
                                                                                                                 Hs.47442
                                                                                                                                                    Hs.93956
                                    Hs.44603
                                                          Hs.14855
                                                                                                                                                                                     Hs.66195
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AA280431
UJ55376
T56281
NA4439
AA089558
NA9500
AA259084
NS2137
NA9537
NA9259
HG2139-
NA457
NA457
NA457
NA457
NA43325
AA40026
T87648
AA400277
R9480
AA400277
R44386
W19088
AA418333
AA43333
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AA4333
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7514	80	AA455914	Hs.1019	Parathyroid hormone receptor 1
866	83	X65644	Hs.75063	HUMAN IMMUNODEFICIENCY VIRUS TYPE I ENHANCER-BINDING PROTEIN 2
962	80	HG180-		EST - HG180-HT180
0540	∞	N62328	Hs.3786	Glutamate receptor metabotropic 3
5700	€	AA131512	Hs.103820	EST
5880	60	AA436706	Hs.98895	ESTs
809	•0	M55267	Hs.41846	EVI2A PROTEIN PRECURSOR TROPIC VIRAL INTEGRATION SITE 2A PROTEIN)
620	e 0	X14885	Hs.2025	Transforming growth factor beta 3
5643	6 0	AA039325	Hs.47200	ESTs
2051	80	R01450	Hs.91061	ESTs
3550	•	W90617	Hs.50120	ESTs
177	80	X75535	Hs.118884	33 KD HOUSEKEEPING PROTEIN
3000	40	W45531	Hs.94642	ESTs
9900	€0	H98701	Hs.4985	ESTs
290	•	U50534	Hs.30649	Human infant brain mRNA clone 13cDNA73
7199	60)	AA448257	Hs.97127	ESTs
4601	89	238844	Hs.25803	ESTs
3589	w	W93074	Hs.59342	ESTs
7389	80	AA453466	Hs.99330	ESTs
1080	80	X55448	Hs,3118	H.sapiens mRNA for 2.19 gene
8748	8	D25912	Hs.74832	ESTs
0000	•	H72592	Hs.77554	ESTs
0367	80	H98857	Hs.75520	Human mRNA for KIAA0041 gene partial cds
3609	80	W93585	Hs.59476	ESTs
4502	8 0	Z38214	Hs.26946	ESTs
1129	€0	AA156873	Hs.15970	ESTs
1144	60	AA033659	Hs.95154	ESTs
8584	€0	C21221	Hs.68619	ESTs Highly similar to METALLOTHIONEIN-IA (Equus caballus)
2165	80	R52822	Hs.22003	ESTs
1987	•0	N94551	Hs.55060	ESTs
2470	e 0	115956	Hs.65289	EST
8642	60	AA599152	Hs.91379	PTB-ASSOCIATED SPLICING FACTOR
0438	€0	N21684	Hs.80500	Human mRNA for KIAA0061 gene partial cds
4471	80	AA258843	Hs.111376	ESTs
1571	Œ	R61005	Hs.115170	Homo saplens mRNA for GaIT4 protein
1417	80	N68435	Hs.49516	ESTs
3951	ø	T97318	Hs.18037	ESTs
7832	4 0	AA249260	Hs.28545	ESTs
7928	&	AA461093	Hs.26799	ESTs Moderately similar to zinc finger protein (R.norvegicus)

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Human DNA sequence from clone 323P24 on chromosome Xp11.21-11.23 Contains SPIN (spindlin homotog (PROTEIN DXF34) hypothetical prote
                                                                                                                                                                                                                                              Human 3'5' cyclic nucleolide phosphodiesterase (HSPDE1A3A) mRNA complete cds
                                                                                                                                                                                                                                                                            Human orphan G-protein coupled receptor Dez isoform a mRNA complete cds
 Celtular retinatidehyde-binding protein
Nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (p105)
                                                                                                                                                                                                                                                          Human Ral guanine nucleotide dissociation stimulator mRNA partial cds
                                                                                                                                                                                                                                                                                                                              Solute carrier family 5 (sodium/glucose cotransporter) member 1
                                                                     ESTs Weakly similar to PROTEIN Q300 [Mus musculus] ESTs
                                                                                                      Human mRNA for KIAA0318 gene partial cds
                                                                                                               EST - RC_AA075674
                                                                                                                                                                                                                                                                                                                                                  Myeloperoxidase
                                                                                                                                                                                                                                                                                                                                                                               EST - X95677
                                                                                                                                   ESTs
EST
ESTs
ESTs
                                                                                                                                                                                                                                                                                                                                                Hs.1817
Hs.124418
Hs.51501
                                                                                                                                                                                                                                   Hs.41717
Hs.41717
Hs.106185
Hs.100739
Hs.111075
Hs.19748
Hs.1933
Hs.83428
Hs.34081
Hs.142935
Hs.33487
                                                 Hs.59736
Hs.31653
Hs.19280
Hs.107894
Hs.103343
Hs.65746
                                                                                                                      Hs.82577
Hs.86723
Hs.29196
Hs.30842
Hs.130843
                                                                                                                                                                         Hs.97699
Hs.49193
Hs.7569
                                                                                                                                                                                                                                                                                               Hs.56213
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                                                                                                                                                                                                                Hs.71016
                                                                                                                                                                                                                                                                                                                             Hs. 1964
                                                                                                                                                                                                                          Hs.55501
                                                                                                                                                                                                      Hs.22057
L34219
AA098834
N55189
AA400155
AA400155
B4400155
H19472
R79356
D60285
AA019426
AA019426
AA039898
R71489
H08171
AA011041
AA398962
N66399
T26893
AA267595
W32094
AA435978
AA40893
U14417
W57731
AA40893
C179527
R06607
W57731
AA40893
R80645
M19507
D80051
D80051
D80051
W63399
R80645
AA283926
R80645
M19507
D80051
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        1952
        1 H0086B
        Hs.31760
        ESTS

        29801
        7 K20339
        Hs.3879
        ESTS

        29807
        7 K20339
        Hs.3879
        ESTS

        23967
        7 K20330
        Hs.3879
        ESTS

        23967
        7 K2030
        Hs.3878
        ESTS

        23967
        7 K2041
        Hs.6018
        ESTS

        2342
        7 K2041
        Hs.6018
        ESTS

        28630
        7 AA28757
        Hs.8024
        ESTS

        28630
        7 AA28767
        Hs.8024
        ESTS

        28630
        7 AA28767
        Hs.8024
        ESTS

        28630
        7 AA28767
        Hs.8021
        ESTS

        7 W8930
        Hs.9421
        ESTS

        7 W8930
        Hs.9421
        ESTS

        7 W8935
        Hs.1042
        ESTS

        7 AA2876
        Hs.1042
        ESTS

        7 W8935
        Hs.1042
        ESTS

        7 AA2876
        Hs.1042
        ESTS

        7 W8935
        Hs.1042
        ESTS

        8 W8935
        Hs.1042
        ESTS

        8 AA2876
        Hs
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ESTs Weakly similar to The KIAA0147 gene product is related to adenylyl cyclase. [H.sapiens]	ESTS TOTAL	EDIS	nomo saprens con e 24323 manas sequence H.sapiens mRNA for ardinne methytransferase	ESTs	ESTs	ESTs	ESTs Weakly similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII [H.sapiens]	ESTs	EST - RC_AA070160	Homo saplens mRNA for KIAA0667 protein partial cds	Human Kox1 gene for zinc finger protein	H.saplens mRNA for melanoma growth regulatory protein MIA	ESTs	ESTs	ESTs	EST	ESTs	EST	ESTs	ESTs Weakly similar to no similarities to reported gene products (H.saplens)	EST Weakly similar to hypothetical protein [H.sapiens]	ESTs	EST - RC_AA070397	ESTs	ESTs	EST	EST	ESTs	Homo sapiens done 24418 mRNA sequence	ESTs	ESTs	Homo sapiens clone 23698 mRNA sequence	ESTs	VON WILLEBRAND FACTOR PRECURSOR	Homo saplens CASK mRNA complete cds	Homo sapiens clone 24700 unknown mRNA partial cds	EST	ESTS
		_	Hs.105365	Hs.119295	Hs.58992	Hs.100119	Hs.19136	Hs.55426		Hs.4217	Hs.104115	Hs.58094	Hs.2906B	Hs.22654	Hs.59872	Hs.50125	Hs.6526	Hs.71057	Hs.22703	Hs.16075	Hs.44792	Hs.65996		Hs.37630	Hs.118898	Hs.67186	Hs.46852	Hs.65373	Hs.13434	Hs.60480	Hs.15345	Hs.8136	Hs.84628	Hs.110802	Hs.35986	Hs.95665	Hs.112509	Hs. 108209
AA609120	AA007591	M68158	N52195	AA442090	W87469	AA130867	AA027317	W23631	AA070160	T10070	X52332	AA282143	AA148983	AA297746	AA002150	N67197	H24317	AA127098	R46597	T87519	N36130	F04014	AA070397	N58628	AA232138	C14820	N48302	T16896	R38804	AA228096	T84046	W25781	W01094	M10321	N92643	C06238	AA599142	W87801
9	v v	o 4	ο φ	9	စ	9	9	9	9	o /	Ф	9	9	9	9	ø	φ	ø	ø	80	w	စ	80	Ø	ø	9	ę,	9	9	9	ø	9	9	9	9	9	9	g
28357	25022	29334	30744	27577	33472	25687	10549	32836	17025	22939	15803	34618	11074	12212	16102	31340	19459	17533	22017	23687	30298	28973	17042	20817	18430	28493	30497	32502	41324	18418	23622	15342	15246	2247	31906	9062	38640	42622

21300	5	10000	20.50	22
7293	g	AA085354		EST - AA085354
27322	φ	AA424325	Hs.40496	ESTs
5774	9	X54199	Hs.82285	Phosphoribosylalycinamide formyltransferase phosphoribosylalycinamide synthetase phosphoribosylaminolmidazole synthetase
33469		W87454	Hs.58987	ESTs Highly similar to homogentisate 12-dioxygenase [H.sapiens]
40329	9	H97488	Hs.108802	Human N-ethylmaleimide-sensitive factor mRNA partial cds
26539	æ	AA278848	Hs.88522	ESTs
23579	ю	T81098	Hs.124065	ESTs
22435	ø	R66706	Hs.28706	ESTs
19956	9	H80842	Hs.37445	ESTs
18607	ø	F02345	Hs.21197	ESTs
18991	စ	H02554	Hs.30323	ESTs
7810	9	AA248169	Hs.106709	ESTs Highly similar to LATENT TRANSFORMING GROWTH FACTOR BETA BINDING PROTEIN 1 PRECURSOR (Ratus novection
30810	0	N53419	Hs.47648	ESTs
17996	ø	AA169606	Hs.72815	ESTs
26245	Ф	AA252357	Hs.87794	ESTs
13348	89	AA449267	Hs.17914	ESTs Weakly similar to HIGH AFFINITY IMMUNOGLOBULIN EPSILON RECEPTOR BETA-SUBUNIT [M.musculus]
15034	•	U43701	Hs.75403	60S RIBOSOMAL PROTEIN L23A
33802	Ф	241058	Hs.79248	ESTs
33394	9	W84432	Hs.58670	ESTs
31170	9	N64017	Hs.48911	ESTs · ·
40828	9	N64144	Hs.102749	EST
1789	9	L13258	Hs.936	Solute carrier family 17 (sodium phosphate) member 2
17740	s n	AA142922	Hs.9817	Homo sapiens Arg/Abi-interacting protein Arg8P2a (Arg8P2a) mRNA complete cds
6266	2	H58970	Hs.49683	ESTs
15540	2	W30895	Hs.7535	ESTs
27827	S	AA455976	Hs.42355	ESTs
30697	ç	N51585	Hs.47049	ESTs
40299	sc.	H95787	Hs.108745	ESTs
16627	ç	AA036779	Hs.61826	Homo sapiens done 23928 mRNA sequence
20659	2	N62915	Hs.57672	ESTs Weakly similar to weakly similar to myosin heavy chain [C.elegans]
31009	2	N62272	Hs.48502	ESTs
14877	40	T65508	Hs.14623	Homo sapiens mRNA for KIAA0625 protein partial cds
31102	ç	N63178	Hs.48728	ESTs
22830	53	R98421		EST - RC_R98421
25248	40	AA040507	Hs.92924	ESTs
32145	ç	R38910	Hs.66170	Homo saplens done 24503 mRNA sequence
27970	2	AA464626	Hs.10247	Activated leucocyte cell adhesion molecule
34509	S	AA262974	Hs.111394	ESTs .
29522	ın	H68338	Hs.90250	ESTs

	ESTs	ESTs	ESTs Weakly similar to kynurenine/alpha-aminoadipate aminotransferase [R.norvegicus]	EST - RC_AA082933	Human mRNA for KIAA0140 gene complete cds	ESTs	Human R kappa B mRNA complete cds	Human hTRIP (hTRIP) mRNA complete cds	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	Human mRNA for BST-1 complete cds	ESTs	ESTs Weakly similar to 3-oxoacyl-{acyl-carrier protein} reductase (E.coli)	ESTs	ESTs	EST - AA412556	Homo sapiens done 23904 mRNA sequence	Sorting nexin 1	_	Myasthenic syndrome antigen B [human fetal brain mRNA 3477 nt]	EST	ESTS	-	EST - D14823	Epidermal growth factor	ESTs	ESTs Weakly simitar to ZINC FINGER PROTEIN 42 [H.sapiens]	ESTs	ESTs	_	Human clone iMAGE:35527 unknown protein mRNA partial cds
	Hs.97505	Hs.99485	Hs.63481		Hs.5084	Hs.70811	Hs.95262	Hs.21254	Hs.7147	Hs.45247	Hs.92262	Hs.48523	Hs.65524	Hs.14651	Hs.9115	Hs.21258	Hs.62264	Hs.37599	Hs.32980	Hs.7508	Hs.94811	Hs.7316	Hs.6166		Hs.67364	Hs.75283	Hs.100543	Hs.30941	Hs.90165	Hs.26236	Hs.101442		Hs.2230	Hs.13251	Hs.1110	Hs.12907	Hs.97600	Hs.136395	Hs.87197
	AA400514	AA459649	AA404282	AA082933	D50930	AA122394	X8087B	U77845	T23513	D80990	R76401	W70259	D45455	R19360	W36290	AA412293	R62579	AA009809	H96306	AA242829	W70279	T23867	W24154	AA412556	C01360	U53225	AA035444	AA039933	D20538	AA096412	AA099580	D14823	X04571	F10040	T62918	T66282	AA398155	N67598	U85992
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ESTs	EST	ESTs	EST	EST - AA249611	ESTs	Homo sapiens mRNA for KIAA0554 protein parital cds	ESTs	EST - RC_H18929	EST - RC_AA016258	ESTs Weakly similar to !!!! ALU SUBFAMILY SO WARNING ENTRY II!! [H.sapiens]	ESTs	ESTs	Laminin alpha 4	ESTs	Human alpha 1-tetoprotein transcription factor (hFTF) mRNA complete cds	Human TNF receptor associated factor 6 (TRAF6) mRNA complete cds	Human LAR-interacting protein 1a mRNA complete cds	ESTs	ESTs	ESTs	STATHMIN	EST	Human DNA damage repair and recombination protein RAD52 pseudogene mRNA partiat cds	ESTs	EST - RC_R05315	ESTs	EST - RC_R01081	ESTS	EST - RC_AA349591	ESTs	ESTs	Human mRNA for KIAA0324 gene partial cds	ESTs	Homo saplens mRNA for KIAA0541 protein partial cds	ESTs	ESTs Highly similar to PUTATIVE 2-HYDROXYACID DEHYDROGENASE IN BISC-CSPA INTERGENIC REGION [Escherichia coif]	ESTs	Phosphatidylinositol glycan class F	ESTs Highly similar to MYELOID DIFFERENTIATION PRIMARY RESPONSE PROTEIN MYD118 (Mus musculus)
Hs.37456	Hs.61425	Hs.16414	Hs.54593		Hs.38323	Hs.74750	Hs.98466			Hs.95278	Hs.55409	Hs.107293	Hs.78672	Hs.117183	Hs.91310	Hs.90957	Hs.5376	Hs.107941	Hs.43616	Hs.55405	Hs.81915	Hs.69293	Hs.73046	Hs.105095		Hs.11923		Hs.55412		Hs.103351	Hs.31181	Hs.7841	Hs.24120	Hs.10881	Hs.99410	Hs.5296	Hs.106385	Hs.111026	Hs.110571
N69215	AA027059	187693	N90168	AA249611	H65459	AA284403	AA426464	H18929	AA016258	AA047078	AA446114	W46403	AA496983	R45175	U93553	078798	H97922	AA156838	AA259058	W20404	D60419	AA101833	U22172	T63336	R05315	AA505141	R01081	W20364	AA349591	AA019598	H12725	AA347209	R26855	AA218543	AA456112	W28944	AA262308	H80865	AA450127
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. 50832	16502	23691	31842	7845	29297	7953	36412	19357	16299	25312	27617	42432	38432	32215	15214	15141	20052	7551	26451	42355	39480	17369	14993	23400	21153	14282	21104	32825	35018	25104	19235	34979	21501	18331	37529	15532	11858	29450	37294

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        33170
        5
        AA3292830
        Hs. 98347
        EST8 Worakly similar to rhp-1 [H. sapiens]

        22011
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        AA389121
        Hs. 9279
        EST8

        22012
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        AA389121
        Hs. 9279
        EST8

        22012
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        AA389121
        Hs. 36718
        Surfacent pulmonary-associated protein D

        20356
        5
        N58009
        Hs. 36718
        EST8
        Worakly similar to hypothetical L1 protein [H. sapiens]

        20376
        5
        N58009
        Hs. 36718
        EST8
        Worakly similar to hypothetical L1 protein [H. sapiens]

        20376
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        N58009
        Hs. 36718
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        Hs. 18009
        Hs. 26018
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27430	4	AA429028	Hs.42676	ESTs Moderately similar to IIII ALU SUBFAMILY SB WARNING ENTRY IIII [H.saplen
30362	4	N40170	Hs.45046	ESTs
30599	4	N50039	Hs.47004	ESTs
27894		AA450319	Hs.48469	ESTs
31844	4	N90218	Hs.54607	ESTs
24935	4	AFFX.		AFFX-HUMGAPDH/M33197_M
40906	•	N68829	Hs.131510	ESTs
35220	4	AA398246	Hs.97594	ESTs
10024	4	N94832	Hs.121699	ESTs
4238	4	U33053	Hs.2499	Human lipid-activated protein kinase PRK1 mRNA complete cds
26358	•	AA256396	Hs.88156	EST
24716	•	Z39734	Hs.22550	ESTs
19243	4	H13539	Hs.31222	ESTs
41844	4	T15833	Hs.100227	EST
39774	4	H17948	Hs.106311	Homo saplens DGS-A mRNA 3' end
20111	4	N21380	Hs.25497	H.saplens mRNA for ROX protein
24008	*	T99337	Hs.18624	ESTs
8316	4	AA410529	Hs.30732	ESTs
35072	4	AA358738		EST - RC_AA358738
39794	4	H21819	Hs.14896	Homo sapiens done 24590 mRNA sequence
1517	4	J04501	Hs.772	Glycogen synthase 1 (muscle)
9164	*	D38081	Hs.89887	Thromboxane A2 receptor
35027	•	AA349996	Hs.96937	ESTs
35185	4	AA398015	Hs.97590	Unitted
26570	*	AA279425	Hs.86428	ESTs Weakly similar to hypothetical L1 protein [H.sapiens]
19759	4	H54352	Hs.36873	ESTs
34310		AA243416	Hs.75470	Homo sapiens mRNA expressed in osteoblast complete cds
39961	4	H57317	Hs.108161	ESTs
23249	4	T47919	Hs.8749	ESTs
19119	•	H09077	Hs.30895	EST
14158	₹	AA490182	Hs.118598	ESTs
22866	4	R99938	Hs.36189	EST
16935	4	AA059392	Hs.66791	ESTs .
41950	4	T33137	Hs.7967	ESTs
20404	-	N48694	Hs.30881	Homo sapiens liprin-alpha2 mRNA complete cds
18406	4	AA227962	Hs.112075	ESTs Weakly similar to reverse transcriptase [M.musculus]
16560	4	AA031308	Hs.24341	ESTs
26098	4	AA242831	Hs.87606	ESTs .
34360	4	AA251547	Hs.104358	EST
46830	7	AA054222	He 40400	

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ESTs Weakly similar to reverse transcriptase homolog [H.sapiens]
                                                                                                                                                                                                                                                                                                                                                                       Human Toll-like receptor 2 (TLR2) mRNA complete cds
                                                                                                                                                                                                                                                                                                                          Human mRNA for KIAA0385 gene complete cds
   Mannose-binding lectin soluble (opsonic defect)
EST - RC_W38051
                                                                                                                                                                                                      Human antigen (MAGE-1) gene complete cds EST - RC_AA121974
                                                                                               EST - AA397529
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R66769
AA398662
AA419995
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AA323299
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KO3474	4 ·			ESTs Highly similar to NUCLEOLYSIN TIA-1 [Homo sapiens]
## R01068 Hs.14603 E ## T16538 Hs.106443 E ## AA406231 Hs.103849 E ## AA406231 Hs.103849 E ## AA406234 Hs.103804 Hs.2536 Hs.103804 Hs.2536 Hs.2643 E ## AA406866 Hs.67337 Hs.09804 Hs.2643 E ## AA4068666 Hs.67337 Hs.09804 E ## AA407612 Hs.60435 E ## AA407612 Hs.60436 E ## AA407613 Hs.51919 E ## AA407614 Hs.51919 E ## AA407615 Hs.106706 E ## AA407616 Hs.6061 Hs.6061 E ## AA407616 Hs.6061 Hs.6061 E ## AA407616 Hs.6061 Hs.106706 E ## AA407616 Hs.6061 Hs.106706 E ## AA407616 Hs.6061 Hs.106706 E ## AA407616 Hs.106706 Hs.106706 E ## AA407616 Hs.106706 Hs.106706 E ## AA407617 Hs.106706 Hs.106706 E ## AA407617 Hs.106706 Hs.106706 E ## AA407617 Hs.106707	,	K03474		EST - K03474
4 AA599661 Hs.103649 EA599661 Hs.103849 EA599661 Hs.103844 EA59969 Hs.2536 Hs.103949 EA59969 Hs.2536 Hs.103949 EA59969 Hs.25399 Hs.25399 Hs.25499 EA59999 Hs.25999	4	R01068	Hs.14603	ESTs
4 AA599661 Hs.103849 E 4 AA466231 Hs.100113 H 4 AA426372 Hs.109804 H 4 AA426372 Hs.109804 H 4 AA4363763 Hs.5366 H 4 AA6057620 Hs.5366 H 4 AA6057620 Hs.536 H 4 AA47612 Hs.60435 H 4 AA47612 Hs.60435 H 4 AA4397616 Hs.60435 H 4 AA4397616 Hs.51919 Hs	4	T16358	Hs. 106443	ESTs
4 AA406231 Hs.100113 H 4 AA26372 Hs.80067 E 4 AA330634 Hs.2336 Hs.2336 Hs.2336 Hs.2336 Hs.2336 Hs.2336 Hs.2336 Hs.2336 Hs.2337 Hs.2007 E 4 AA4057620 Hs.2007 E 4 AA4057620 Hs.2007 E 4 AA4057620 Hs.2007 E 4 AA407612 Hs.60435 Hs.2137 Hs.2130 Hs.21316 Hs.20136 Hs.21316 Hs.20136 Hs.2014 Hs.2016 Hs.2016 Hs.2016 Hs.2016 Hs.2016 Hs.2016 Hs.2016 Hs.2016 Hs.2013 Hs.2016 Hs.2013 Hs.20	4	AA599661	Hs.103849	ESTs
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4 AAA26372 Hs.109804 H 4 AAA330534 Hs.2536 H 4 NB984 Hs.5536 Hs.0537 Hs.2643 Hs.67317 Hs.0447612 Hs.0435 Hs.106435 Hs.107245 Hs.2514 Hs.0435 Hs.107245 Hs.2614 Hs.660435 Hs.2614 Hs.	4	D20261	Hs.80067	ESTs
4 AA330634 4 ABS2R8 H8.2536 H8.08984 H8.64543 H8.64543 H8.6451317 H8.6451317 H8.64513 H8.6601 H8.64513 H8.6601 H8.64513 H8.6601 H8.64513 H8.6601 H8.	4	AA426372	Hs.109804	Human mRNA for histone H1x complete cds
235278 Hs.2536 Hs.0536 Hs.0536 Hs.0536 Hs.053717 Hs.043717 Hs.043717 Hs.043717 Hs.043717 Hs.043717 Hs.043717 Hs.043717 Hs.043717 Hs.043717 Hs.0437 Hs.2514 Hs.0514 Hs.	₹.	AA330634		EST - RC_AA330634
AA057620 Hs.54543 RA057620 Hs.50807 RA0689696 Hs.6717 RA047612 Hs.60436 Hs.6177 RA047612 Hs.60436 Hs.61774 RA047612 Hs.60436	•	Z35278	Hs.2536	H.sapiens PEBP2aC1 acute myeloid teukaemia mRNA
4 AA057620 H9.30807 RA068666 H9.67317 RA068666 H9.67317 RA26844 R9.67317 R9.6944 R9.69	4	N89848	Hs.54543	ESTs
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4 AA283393 Hs.87734 AA287097 Hs.20714 D88152 Hs.507195 Hs.57191 AA484715 Hs.6000 Hs.20706 Hs.2160 Hs.21613	4	AA447612	Hs.60435	ESTs
4 AA397616 Hs.107245 5 AA287097 Hs.25114 6 NS2979 Hs.25114 7 AA444115 Hs.6000 7 AA48715 Hs.2160 7 AA487165 Hs.105706 7 AA487165 Hs.105706 7 AA487169 Hs.60339 7 AA243172 Hs.60339 7 AA447759 Hs.120969 7 AA47759 Hs.120969 7 AA47759 Hs.120969	4	AA253393	Hs.87734	ESTS
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NB9155	4	AA287097	Hs.25114	ESTs
4 N52979 Hs.51919 4 AA454115 Hs.6000 4 H94266 Hs.105706 4 AA487165 Hs.083008 4 AA442669 Hs.68061 5 AA422646 Hs.68061 4 AA422669 Hs.68051 5 AA423669 Hs.68051 6 AA423669 Hs.680359 7 AA424759 Hs.87619 8 SR2769 Hs.104133 9 AA244172 Hs.120969 3 AA447759 Hs.134724 3 AA447759 Hs.134724 3 AA447159 Hs.22515 3 AA014877 Hs.22515 3 AA014897 Hs.225137	~	D88155	Hs.97196	Human steroidogenic factor 1 mRNA complete cds
4 AA454115 Hs.6000 4 N68730 Hs.12160 4 R95778 Hs.33008 4 AA442669 4 AA442669 4 AA442669 4 AA42669 4 AA42669 6 AA42669 7 AA44269 7 AA44269 7 AA44269 7 AA44269 7 AA44269 7 AA44269 7 AA4269 7 AA44269 7 AA44269 7 AA447759 Hs.120969 7 AA447759 Hs.134724 7 AA47759 Hs.25515 7 AA471360 Hs.25515 7 AA674897	4	N52979	Hs.51919	Plasmhogen-like protein
N68730 H5.12160 H94266 H5.9451 H94266 H5.9451 H859778 H5.105706 H8442669 H8442669 H8442669 H846061 H8460	4	AA454115	Hs.6000	ESTs
4 H94266 Hs.9451 4 R65778 Hs.3008 4 AA44765 Hs.105706 4 AA422669 Hs.68061 4 AA013070 Hs.68033 4 AA010070 Hs.89359 5 S62769 Hs.104133 3 AA243172 Hs.87619 3 W92001 Hs.120969 3 AA447759 Hs.13245 3 AA447759 Hs.13265 3 AA447769 Hs.22515 3 AA014897 Hs.22515 3 AA014897 Hs.22517	4	N68730	Hs.12160	ESTs
4 R95778 H5.33008 4 AA487165 H5.105706 4 AA442689 4 AA232646 H5.68061 4 AA010070 H5.60339 4 W49755 H5.89359 3 AA243172 H5.897619 3 AA24172 H5.87619 3 AA24175 H5.87619 3 AA24176 H5.120969 3 AA241769 H5.120969 3 AA212160 H5.22615	4	H94266	Hs.9451	ESTs
4 AA487165 Hs.105706 4 AA422669 4 AA6232646 Hs.68061 4 AA010070 Hs.60339 4 W49755 Hs.89359 4 W49755 Hs.89359 3 AA243172 Hs.87619 3 W92001 Hs.120969 3 W92001 Hs.120969 3 AA447759 Hs.134724 3 AA41736 Hs.2515 3 AA074897 3 AA074897 3 F04262 Hs.22137	4	R95778	Hs.93008	EST
AA442669 A AA222646 Hs.68061 A AA010070 Hs.60339 A W49755 Hs.104133 AA243172 Hs.87619 A W92001 Hs.120969 A AA47759 Hs.134724 AA447759 Hs.73724 AA014897 AA014887 AA014887 AA014887 AA014887	4	AA487165	Hs.105706	EST
4 AA232646 Hs.68061 4 AA010070 Hs.60339 4 S82769 Hs.104133 3 AA234172 Hs.87619 3 W92001 Hs.120969 3 AA47759 Hs.134724 3 AA014897 3 AA014897 3 F04262 Hs.22137	4	AA442669		EST - AA42669
4 AA010070 Hs.60339 4 W49755 Hs.69359 4 S87769 Hs.104133 3 AA243172 Hs.87619 3 W92001 Hs.120969 3 AA447759 Hs.13069 3 AA47759 Hs.137567 3 AA074697 3 AA074697 3 F04262 Hs.22137	4	AA232646	Hs.68061	ESTs
4 W49755 Hs.69359 4 S82769 Hs.104133 3 AA243172 Hs.87619 3 W92001 Hs.120969 3 AA447759 Hs.134724 3 H10047 Hs.22515 3 AA4121360 Hs.27567 3 AA074897 3 F04262 Hs.22137	4	AA010070	Hs.60339	EST
\$62769 Hs.104133 \$AA243172 Hs.87619 \$W92001 Hs.120969 \$A447759 Hs.134724 \$A447759 Hs.134724 \$AA47759 Hs.22515 \$AA074897 \$AA074897 \$AA074897 \$AA074897	~	W49755	Hs.89359	ESTs Moderatety similar to type la hair keratin a3 (H.sapiens)
3 AA243172 Hs.87619 3 W92001 Hs.120969 3 AA447759 Hs.134724 3 AA014897 3 AA074897 3 F04262 Hs.22515	•	\$82769	Hs.104133	GABAA receptor garrma 3 subunit [human fetat brain mRNA Partial 1536 nt]
3 W92001 Hs.120969 Hs.134724 Hs.134724 Hs.134724 Hs.22515 Hs.22515 Hs.22515 Hs.27567 Hs.27567 Hs.27567 Hs.22137	6	. AA243172	Hs.87619	ESTs
3 AA447759 Hs.134724 13 H10047 Hs.22515 3 AA121360 Hs.27567 3 AA074897 3 F04262 Hs.22137 1	က	W92001	Hs.120969	ESTs
3 H10047 Hs.22515 3 AA121360 Hs.27567 3 AA074897 3 F04262 Hs.22137	c,	AA447759	Hs.134724	ESTs
3 AA121360 Hs.27567 13 AA074897 3 F04262 Hs.22137	٣	H10047	Hs.22515	ESTs
3 AA074897 6 3 F04262 Hs.22137 6	ы	AA121360	Hs.27567	ESTs Weakly similar to similar to alpha-13-mannosyl-glycoprotein beta-1 2-N-acetylgtucosaminyltransferase (C.etega
3 F04262 Hs.22137	e,	AA074897		EST - AA074897
	က	F04262	Hs.22137	ESTs
3 N77904	e	N77904	Hs.44380	ESTS .
40997		• • • • • • • • • • • • • • • • • • • •	AA599661 AA406231 AA406231 AA406231 AA426372 AA330634 AA330634 AA4069696 AA065696 AA065696 AA065696 AA065696 AA065696 AA065696 AA065696 AA065696 AA065696 AA06399 AA330634 AA44269	

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ESTs	Homo sapiens mKNA Ior osteoblast specific cysteine-rich protein complete cos		PROTEIN KINASE CITHETA TYPE	EST - RC_H28966	5-HYDROXYTRYPTAMINE 28 RECEPTOR	ESTs Weakly similar to ETX1 (alternatively spliced) [H.sapiens]	Transforming growth factor alpha	ESTs .	ESTs	ESTs Highly similar to HYPOTHETICAL 70.7 KD PROTEIN F09C8.3 IN CHROMOSOME III (Caenomabdikis elegans)	ESTs	ESTs ·	Homo sapiens done 23837 mRNA sequence	Homo sapiens done 24466 mRNA sequence	ESTs	EST	Inositol polyphosphate phosphatase-like protein 1 (51C protein)	ESTs	ESTs .	ESTs	ESTs	ESTs	Human GT334 protein (GT334) gene mRNA complete cds	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	N-acetylglucosaminyitransferase I	ESTs	ESTS	ESTs	ESTs	EST
Hs.62440	HS.62562	18.23351	Hs.89615		Hs.2507	Hs.23153	Hs.2023	Hs.13759	Hs.4248	Hs.30490	Hs.23336	Hs.98124	Hs.110480	Hs.25924	Hs.23539	Hs.141719	Hs.75339	Hs.23352	Hs.12345	Hs.106291	Hs.99566	Hs.22928	Hs.94479	Hs.138717	Hs.93090	Hs.27109	Hs.72163	Hs.35088	Hs.14633	Hs.22660	Hs.31697	Hs.101504	Hs.117946	Hs.74876	Hs.11590	Hs.40478	Hs.110964	Hs.68513
AA180487	AAU47265	450034	R16896	H28966	N36174	R33005	X70340	170580	AA412620	N48329	R22057	AA412290	AA283907	H10068	Z41301	W68846	L36818	W23474	F10565	C02049	AA454935	H10641	AA303078	N49952	AA398488	F03004	AA157291	W07019	N51599	AA142849	H23747	R69233	M55621	D20188	AA478441	H83694	AA151621	C20680
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11217	74/37	11417	21382	19508	30301	21553	6102	23502	6333	30500	21431	35920	12065	19156	24844	33178	2052	15327	18874	9039	37470	19167	34888	30591	26997	18647	17867	15280	20465	11047	19451	41621	2822	28675	13928	29473	25829	28532

32370	, ,	181381	HS.04393	8704
12064	es.	AA283848	Hs.11367	ESTs Weakly similar to KIAA0009 [H.sapiens]
15547	6 0	W32012	Hs.29353	ESTs
40284	က	H95073	Hs. 108734	ESTs Highly similar to HYPOTHETICAL 55.1 KD PROTEIN IN FAB1-PES4 INTERGENIC REGION (Saccharomyces cerevisiae)
22861	e	R99595	Hs.36152	ESTs
15525	9	W28763	Hs.16349	Homo sapiens KiAA0431 mRNA partial cds
17968	en	AA167496	Hs.72695	EST
3836	ဗ	U12707	Hs.2157	Wiskott-Aldrich syndrome (ecezema-thrombocytopenia)
4528	က	U48251	Hs.75871	Homo sapiens protein kinase C-binding protein RACK7 mRNA partial cds
29784	т.	N20468	Hs.42849	ESTs Weakly similar to line-1 protein ORF2 [H.sapiens]
7892	က	AA262100		EST - AA262100
15279	6	W05746	Hs.133302	ESTs
42064	e	T63364	Hs.9225	ESTs Highly similar to FIBRINOGEN-LIKE PROTEIN A PRECURSOR (Parastichonis nandmensis)
17943	ო	AA165117	Hs.20509	ESTS
4596	ET.	U50743	Hs.19520	Sodiumpotassium ATPase gamma subunit
16421	60	AA022541	Hs.61146	ESTs
4914	e	U67611		EST - U67611
38171	6	AA487301	Hs.105713	EST
20168	es	N24106	Hs.2799	Cartilage linking protein 1
18791	ъ	F09892	Hs.12575	ESTs
24281	n	W79773	Hs.16511	ESTs
13996	ო	AA480907	Hs.15769	ESTs
20583	ო	N57797	Hs.34421	ESTs
37181	n	AA448158	Hs.99152	EST
21450	۳	R23146	Hs.23466	ESTs
19634	e	H44866	Hs.31597	ES1s
10163	က	R54534	Hs.7898	Homo sapiens done 23938 mRNA sequence
7059	က	296810		EST - 296810
25762	က	- AA136068	Hs.91797	Retinoblastoma-binding protein 1 (alternative products)
27426	e	AA428900	Hs.92897	ESTs
29023	e	F09302		EST - RC_F09302
10989	ო	AA132366	Hs.8023	Homo sapiens mRNA for SPOP
6587	8	X97748		EST - X97748
8722	3	AA481309	Hs.30204	ESTs
21476	۳	R26065	Hs.23523	ESTs
14096	es!	AA487558	Hs.8135	ESTs
35392	m	AA399562	Hs.97566	ESTs
28608	က	C21509	Hs.112774	ESTs
13350	es	AA449297	Hs.8944	ESTs Weakly simitar to type 1 procollagen C-proteinase enhancer protein precursor [H.sapiens]
41202	ъ	R12808	Hs.113619	ESTs

A A A 23865 H: 1052 UTROPHIN A H: 10645 ESTS Weakly similar to coded for by C. elegans CDNA cm10e3 [C.elegans] H: 1072 H: 10645 ESTS Weakly similar to coded for by C. elegans CDNA cm10e3 [C.elegans] H: 10645 ESTS Weakly similar to C1-TETRAH/CROPCOLATE SYNTHASE CYTOPLASMIC [Rattus norvegicus] H: 10630 H: 10630 ESTS H: 10645 ESTS Highly similar to C1-TETRAH/CROPCOLATE SYNTHETASE CYTOPLASMIC [Rattus norvegicus] A A A 44253 H: 10645 ESTS A A A 44253 H: 10645 ESTS A A 46254 H: 10645 ESTS A 46254 H: 10645 ESTS A 46254 H: 10645 ESTS ESTS ESTS ESTS ESTS ESTS ESTS EST
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Hs. 73372 Hs. 92820 Hs. 110454 Hs. 26750 Hs. 110454 Hs. 1482 Hs. 1482 Hs. 1482 Hs. 1674 Hs. 1674 Hs. 1674 Hs. 1674 Hs. 1674 Hs. 23161 Hs. 23161 Hs. 23161 Hs. 23161 Hs. 23037 Hs. 23
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Hs. 73372 Hs. 95870 Hs. 104252 Hs. 110454 Hs. 26750 Hs. 100383
HS.133/2 HS.95870 HS.104252 HS.110454 HS.26750
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Hs.95870 F Hs.95870 F Hs.104252 (
Hs./33/2 Hs.95870
HS./33/2

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ESTs Highly similar to HYPOTHETICAL 54.9 KD PROTEIN C02F5.7 IN CHROMOSOME III (Caenorhabditis elegans)
                        ESTs Highly similar to OPIOID BINDING PROTEIN/CELL ADHESION MOLECULE PRECURSOR [Bos taurus]
 ESTs Weakly similar to The ha1237 gene product is related to S.pombe rad21 gene product. [H.sapiens]
             ESTs Weakly similar to The KIAA0147 gene product is related to adenylyl cyclase. [H.sapiens]
                                                              Human brain mRNA homologous to 3'UTR of human CD24 gene partial sequence
                                      Homo sapiens heat shock factor binding protein 1 HSBP1 mRNA complete cds
                                                                                                                                                                ESTs Moderately similar to sodium-calcium exchanger form 3 [R.norvegicus]
                                                                                                                                                                                                                                                                                                       Homo sapiens retinoic acid-inducible endogenous retroviral DNA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   IMMUNOGLOBULIN-RELATED 14.1 PROTEIN PRECURSOR
                                                                                                                                                                                        Human mRNA for KIAA0008 gene complete cds
Acyf-Coenzyme A dehydrogenase C-2 to C-3 short chain
                                                                                                                                                                                                                                                      Probable transcription factor PML (alternative products)
                                                                                                                                                                                                                                                                                                                                                                                                        Human LIM protein MLP mRNA complete cds
                                                                                                                                                                                                                                                                              Homo saplens done 23872 mRNA sequence
                                                                                                                                                                                                                                                                                                                                                                                Homo saplens drp1 mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Nitric oxide synthase 3 (endothellal cell)
                                                                                                                                                                                                                              EST - U64573
                                                                                                              ESTs
                                                                                                                                     Hs.131828
                                               Hs.125146
Hs.83525
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Hs.23748
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N01031
H02255
R92181
T41177
F01360
N91968
N91968
M4499911
M51335
M44232206
D20378
M92734
M463858
M4403446
M97734
M463496
AA496891
AA131394
Z38804
                                               AA300151
AA167051
W86007
                                                                                                 Z338522
R61750
N34288
R67468
R52145
AA021284
H66642
Z21217
M26393
N72295
                                    A095885
                                                                                      R79580
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EST	ESTs	ESTs	ESTs	ESTs	Protein kinase C substrate 80K-H	Flavin-containing monooxygenase 4	ESTs	EST	ESTs Weakly similar to No definition line found [C.elegans]	CDW52 antigen (CAMPATH-1 antigen)	ESTs	Homo saptens mRNA for SH3 binding protein complete cds done; RES4-23A	ESTs Weakty similar to C06G8.3 [C.elegans]	ESTs Moderately similar to retinoid X receptor interacting protein [M.musculus]	EST - RC_AA063316	ESTs	Homo saplens bicaudal-D (BICD) mRNA complete cds	ESTs	EST - RC_T97353	EST - RC_N45221	Homo sapiens DBI-related protein mRNA complete cds	EST - T47519	ESTs Moderately similar to located at OATL1 [H.sapiens]	Phosphodiesterase 8A cGMP-specific rod alpha	ESTs	ESTs	ESTs Weakly similar to F59C6.4 [C.elegans]	ESTs .	ESTs	EST .	ESTs Weakly similar to ORF YDL077c [S.cerevisiae]	EST - HG1804-HT1829	ESTs	ESTs	ESTs	ESTs Weakly similar to hypothetical protein [H.sapiens]	EST - RC_AA121338	ESTs Weakly similar to reverse transcriptase homolog (H.sapiens)	ESTS
Hs.94667	Hs.14632	Hs.112575	Hs.12296	Hs.75429	Hs.1432	Hs.2664	Hs.142528	Hs.48058	Hs.37477	Hs.108338	Hs.7973	Hs.16227	Hs.5260	Hs.11797		Hs.104747	Hs.24912	Hs.60669			Hs.15250		Hs.98220	Hs.63260	Hs.133217	Hs.20231	Hs.64147	Hs.91681	Hs.89121	Hs.95875	Hs.9452		Hs.32060	Hs.109968	Hs.29696	Hs.10175		Hs.65826	Hs.104186
W46976	N75055	AA608577	N48293	R87373	R63695	Z11737	T86826	N57730	N73988	X62466	H16976	AB000463	R49689	239406	AA063316	AA406219	AA464267	AA058659	T97353	N45221	W79046	T47519	AA417344	W28798	R84933	R08773	AA234687	N74336	AA284722	AA156504	AA043115	HG1804-	AA401452	H41235	W70158	N93764	AA121338	F03032	AA215637
м	7	2	7	7	~	7	7	7	2	7	2	7	7	2	7	2	2	7	7	~	~	~	~	~	7	7	7	7	~	7	~	7	2	~	7	~	7	7	~
33022	31704	38713	20396	10310	22388	15936	23667	30903	20938	5935	19304	8804	41485	24685	25403	35773	27965	16911	42315	40632	15722	14842	36088	15527	10302	21243	7759	31672	26799	25857	16695	96	12439	19599	24223	21052	17463	28949	34140

EST - AA120886	ES) - I'VXVIUSTA GONE 2440 mRNA sequence	Homo sapiens clone 24525 mRNA sequence	ESTs	EST - AA359093	ESTs	ESTS	ESTS	ESTs	ESTs	ESIS	Glutathione S-transferase M5	ESTs	ESTs	ESTs Weakly similar to KIAA0412 [H.sapiens]	EST - HG1019-HT1019	EST - RC_T91086	ESTs	ESTS	Human RACH1 (RACH1) mRNA complete cds	ESTs	ESTs	ESTs	ESTs	Human RNA-binding protein CUG-BP/hNab50 (NAB50) mRNA complete cds	ESTs	Zinc finger protein 136 (clone pHZ-20)	V-ski avian sarcoma viral oncogene homolog	Human Hpast (HPAST) mRNA complete cds	Human clone ID 193225 NAD (H)-specific isocitrate dehydrogenase gamma subunit mRNA atternatively spliced partial cds	ESTS Highly similar to PROTEIN PHOSPHATASE PP2A 55 KD REGULATORY SUBUNIT NEURONAL ISOFORM (Oryctolagus cuniculus)	ESTs	ESTs	ESTS	ESTs	ESTs	ESTS	EST	EST - W/76399
	Hs.85053	Hs.132744	Hs.143798		Hs.22482	Hs.13269	Hs.3757	Hs.42519	Hs.107680	Hs.10444	Hs.75652	Hs.110837	Hs.132188	Hs.18995			Hs.88550	Hs. 104900	Hs.64607	Hs.23710	Hs.70724	Hs.54960	Hs.49759	Hs.81248	Hs.112944	Hs.69740	Hs.81972	Hs.7214	Hs.75253	Hs.108081	Hs.117619	Hs.124027	Hs.57911	Hs.20755	Hs.10024	Hs.85564	Hs.31040	
AA120886	U79288	-	H48488	AA359093	H11509	N50785	AA011310	H98244	H37909	C00185	R40442	AA436156	W88550	F03989	HG1019-	T91086	AA279089	AA453381	N47688	R45441	AA120766	N93495	N69850	T16389	AA621067	R34073	Y09846	AA281769	H18412	H50178	H09751	AA424179	AA025903	R11208	AA176446	AA187955	H11274	W76399
~ ~	, 7	8	7	2	7	81	7	,	7	7	2	7	7	61	7	~	8	8	7	7	2	7	2	7	7	2	7	7	7	7	7	7	2	7	7	2	7	7
7465	5130	30041	19684	8166	19202	20439	10431	29707	39868	8988	41350	13121	15747	18674	914	23804	26556	8567	30457	21975	17452	31958	31495	32490	39174	21572	15914	12014	39777	9484	19147	36276	16475	21304	11189	18093	19190	15710

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Growth hormone 1
ESTs Moderately similar to !!!! ALU CLASS B WARNING ENTRY !!!! [H.sapiens]
                                                                             EXTRACELLULAR SIGNAL-REGULATED KINASE 3
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Homo saplens done 24510 mRNA sequence
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Human RGP3 mRNA complete cds
ESTs Weakly similar to D2045.9 [C.elegans]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Homo sapiens clone 23718 mRNA sequence
                              ranscription factor COUP 2 (a.k.a. ARP1)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Peroxisomal biogenesis factor 6
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               DNA-BINDING PROTEIN A
                                                                                                                                                                                                                                                                                                                                                                                                                                 ST - RC_AA084723
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 EST - RC_AA129060
                                                                                                                                                                                                                                                                                                    EST - RC_H12243
                                                                         Hs. 75649
Hs. 35104
Hs. 19066
Hs. 109047
Hs. 712272
Hs. 7765
Hs. 26054
Hs. 22552
Hs. 23805
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Hs.10688
Hs.115352
Hs.97283
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Hs.89491
Hs.86641
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Hs.23954
Hs.59718
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Hs.19978
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F04627
X91504
N68669
AA018601
A441988
AA42130
AA42130
AA447988
AA42889
T47601
H12243
T47601
H12243
AA428989
T47601
C6072
J03071
AA28305
AA084723
AA084723
AA084723
AA084723
AA287651
AA084716
N24730
N29866
N57007
AA480045
N57007
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  18707

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37400	7	AA453578	Hs.120994	ESTs Weakly similar to T20D3.5 [C.elegans]
29487	2	H85120	Hs.80881	N-ACETYLLACTOSAMINE SYNTHASE
10197	7	R64199	Hs.50785	Homo sapiens vesicle trafficking protein sec22b mRNA complete cds
322	7	D21241		EST - D21241_xpt1
22224	7	R55192	Hs.25689	ESTs
19488	~	H27675	Hs.25604	ESTs
10568	7	AA029703	Hs.36574	ESTs
10872	7	AA099357	Hs.15780	ESTs Moderately similar to ATP-BINDING CASSETTE TRANSPORTER 2 [Mus musculus]
29987	2	N26172	Hs.43760	ESTs
30799	7	N53143	Hs.64001	ESTs
9638	7	L07592	Hs. 106415	Human peroxisome proliferator activated receptor mRNA complete cds
21069	7	N98461	Hs.17706	ESTs
27195	2	AA411473	Hs.65311	ESTs
9241	7	D79565		EST - D79565
9218	2	D61469	Hs.56896	ESTs
22238	7	R55763	Hs.107287	ESTs
7512	8	AA136369		EST - AA136369
17438	2	AA115508	Hs.2780	Jun D proto-oncogene
24932	7	AFFX-		AFFX-HSAC07/X00351_M
28911	8	F01525	Hs.3786	Glutamate receptor metabotropic 3
10944	2	AA125969	Hs.34769	ESTs Weakly similar to F35G12.9 (C.elegans)
42324	~	198199	Hs.142702	ESTs
34756	2	AA287665	Hs.8245	ESTs
12743	2	AA421050	Hs.24545	ESTs
21275	7	R10075	Hs.14890	ESTs
13676	2	AA459389	Hs.26350	Homo sapiens mRNA for tyrosyl sulfotransferase-2
13009	2	AA430474	Hs.16466	ESTs
7403	7	AA094921	Hs.79788	ESTs
11151	7	AA159961	Hs.25819	ESTs
35669	~	AA404707	Hs.54865	ESTs
17701	7	AA135941	Hs.71626	ESTs
18713	2	F04686	Hs.21782	ESTs
24144	7	W52312	Hs.30303	ESTs Highly similar to ZINC FINGER PROTEIN 45 [Homo sapiens]
30741	7	N52175	Hs.22222	ESTs
23042	7	T23433	Hs.7105	ESTs
18479	7	AA233483	Hs.87159	ESTs
8314	7	AA410355	Hs.103081	ESTs Highly similar to RIBOSOMAL PROTEIN S6 KINASE [Homo saplens]
7990	7	AA291786	Hs.32822	ESTs Weakly similar to HYPOTHETICAL PROTEIN KIAA0168 [H.sapiens]
10388	2	AA005355		EST - RC_AA005355

2407	,	HOODBE		EST. HYMBE
42791	. ~	AFFX-		AFEX-HIMCAPDHMA3197 M
26302	8	AA255470	Hs.88040	ESTS
30722	-	N51924	Hs.47370	ESTs
24965	-	AFFX.		AFFX-LysX-5
38850	-	AA609262		EST - RC_AA609262
13746	-	AA461300	Hs.30643	ESTs
6893	-	Z30643	Hs.123123	H.saplens mRNA for chloride channel (putative) 2139bp
31403	-	N68117	Hs.41119	ESTs
17830	-	AA152312	Hs.72047.	ESTs
10583	-	AA033874	Hs.12035	ESTs
20913	-	N72116	Hs.57435	Natural resistance associated macrophage protein 2
. 35607	-	AA402267	Hs.133475	ESTs Weakly similar to zinc finger protein [H.sapiens]
9920	-	N44756	Hs.141935	ESTs Weakly similar to transformation-related protein [H.saplens]
9468	-	H46074	Hs.31562	ESTs
29469	-	H82929		EST - RC_H82929
18121	-	AA191310	Hs.89608	Protein phosphatase 2 (formerty 2A) regulatory subunit A (PR 65) beta isoform
14705	-	D59362		EST - RC_D59362
18692	-	F0444	Hs.6217	ESTs /
16543	-	AA029430	Hs.61557	EST
16359	-	AA019197	Hs.40763	ESTs
40818	-	N63772	Hs.144550	Homo sapiens DNA sequence from PAC 434014 on chromosome 1q32.3.41. Contains the HSD11B1 gene for Hydroxysteroid (11-beta)
35205	-	AA398161	Hs.97602	ESTs
7831	-	AA249175	Hs.96334	ESTs Weakly similar to GOLIATH PROTEIN [Drosophila melanogaster]
. 46771	-	AA150260	Hs.72062	EST
11347	-	AA224245	Hs.26612	ESTs ·
17919	-	AA161125	Hs.72384	EST
22184	-	R53520	Hs.102755	ESTs
14827	-	T35288	Hs.90421	ESTs
28091	-	AA481788	Hs.87593	ESTs
28815	-	D59267	Hs.58782	ESTs
17813	-	AA151480	Hs.91202	ESTs
24655	-	239191	Hs.27262	ESTs Weakly similar to Lph17p [S.corevisiae]
15611	-	W51743	Hs.35096	ESTs
39998	-	H62865	Hs.38132	ESTs
15700	-	W73859	Hs.78061	Homo saplens mesoderm-specific basic-helix-loop-helix protein (FOD1) mRNA complete cds
22045		R48965	Hs.24796	EST
36770	-	AA435753		EST - RC_AA435753
2486	-	N31127	Hs.40098	ESTs
22467	-	R68284	Hs 28399	ESTS

32400		R97176 AA069425	Hs.110783 Hs.20573	ESTs ESTs
23033		T17353	Hs. 108894	Homo sapiens done 23918 mRNA sequence
17593	-	AA129856		EST - RC_AA129856
14867	-	T58588	Hs.100419	ESTs
20266	-	N32118	Hs.107365	ESTs Highly similar to THREONYL-TRNA SYNTHETASE CYTOPLASMIC [Homo sapiens]
14447	-	AA609045	Hs.11759	ESTs
37994	-	AA479919		EST - RC_AA479919
12892	-	AA426521	Hs.18528	Homo sapiens mRNA for NA14 protein
11970		AA280587	Hs.4069	ESTs
19738	-	H53059	Hs.15548	ESTs
14471	-	AA609346	Hs.20102	ESTs
14855	-	154762	Hs.9786	ESTs
24725	-	Z39781	Hs.8004	. Homo sapiens Duo mRNA complete cds
9625	-	X55019	Hs.99975	Cholinergic receptor nicotinic delta polypeptide
20259	-	N31598	Hs.12727	ESTs
18441	-	AA232508	Hs.77480	ESTs
18468	-	AA233177	Hs.87134	ESTs
10164	-	R54743	Hs.19400	ESTs Weakly similar to MITOTIC MAD2 PROTEIN (S.cerevisiae)
8830	-	AB002319	Hs.8663	Human mRNA for KIAA0321 gene partial cds
15287	-	W07461	Hs.121559	ESTs Highly similar to DIPHTHINE SYNTHASE (Saccharomyces cerevisiae)
16477	-	AA026031	Hs.61312	ESTs
21969	-	R45334	Hs.21175	ESTs
22340	-	R61522	Hs.26892	ESTs
12884	-	AA426259		EST - RC_AA426259
8682	-	AA477891	Hs.104476	ESTs
22594	-	R79793	Hs.29900	ESTs
19131	-	H09331	Hs.122791	ESTs
17103	-	AA074997		EST - RC_AA074997
35620	-	AA402493	Hs.10552	ESTs Highly similar to c-Jun feucine zipper interactive [M.muscutus]
35401	-	AA399593	Hs.97682	EST
10901	-	AA112307	Hs.25224	ESTs
19546		H37901	Hs.32706	ESTs
15378	-	W26448	Hs.15071	ESTs Highly similar to VACUOLAR ATP SYNTHASE SUBUNIT D [Bos taurus]
30292	-	N35978	Hs.82364	ESTs
39087	-	AA620607	Hs.111591	ESTs
37896	-	AA477463	Hs.77039	Ribosomal protein S28
8836	-	AB002325	Hs.105917	Human mRNA for KIAA0327 protein complete cds
16150	-	AA005428	Hs.60140	ESTs
23955	-	197467	Hs. 18065	ESTs Weakly similar to alternatively spliced product using exon 13A [H.sapiens]

7812	~	AA469939	Hs.105323	ESTs
4782		S72370	Hs.89890	Pyruvate carboxylase
3540	-	179178	Hs.14463	ESTs
1552	-	R59352	Hs.101253	
1836	-	R42569	Hs.22444	EST
1467	-	AA234089	Hs.14593	ESTs
8347	-	AA219230	Hs.86815	ESTs
215	-	AA389673	148.84344	ESTs Weakly similar to No definition line found [C.elegans]
5205	-	W28366	Hs.7252	Homo sapiens done 24800 mRNA sequence
2764	-	R93802	Hs.33687	ESTs
4966	-	007620	Hs.89661	Human MAP kinase mRNA complete cds
4213	-	W69184	Hs.23973	ESTs
165	-	AA35888	Hs.104135	Homo sapiens mRNA for DRIM protein
2724	-	T90750	Hs.15230	ESTs Weakly similar to F25H2.2 [C.elegans]
212	-	AA386236	Hs.96757	Homo saplens transcription factor SUPT3H (SUPT3H) mRNA complete cds
834	-	M92299	Hs.22554	Homeo box B5 (2.1 protein)
229	-	AA059213	Hs.91898	ESTs
5649	-	W58725	Hs.75074	H.saplens niRNA for MAP kinase activated protein kinase
2306	-	T96538	Hs.45090	Human K+ channel beta 1a subunit mRNA alternatively spliced complete cds
159	-	D31483	Hs.90062	Homo saplens clone 23565 unknown mRNA partial cds
0040	-	H96712	Hs.125198	
2218	-	T86444	Hs.110095	ESTs
5526	-	W28790	Hs.8124	ESTs
7790	-	AA150182	Hs.42262	ESTs Weakly similar to HYPOTHETICAL 88.1 KD PROTEIN K02D10.1 IN CHROMOSOME JII [C.elega
7595	-	AA129929	Hs.71166	EST
1314	-	N66866	Hs.49278	EST
111	-	M57888	Hs.95946	GRANZYME H PRECURSOR
5373	-	W26376	Hs.74563	THY-1 MEMBRANE GLYCOPROTEIN PRECURSOR
3547	-	179448	Hs.14577	EST
2076	-	AA284362	Hs.6448	ESTs Weakly similar to No definition line found (C.elegans)
1956	-	AA279991	Hs.124691	ESTs
5391	-	W26651	Hs. 15961	ESTs
287	-	082557	Hs.18104	ESTs
6419	-	AA022466	Hs.61141	EST
1713	-	R39930	Hs.21034	ESTs
2905	-	AA427537	Hs.32419	ESTs
0257	-	N34961	Hs.75873	H.sapiens mRNA for Zyxin
8134	-	AA487622	Hs.64095	ESTs
6380	-	AA019750	Hs.40797	ESTs Weakly similar to LINE-1 REVERSE TRANSCRIPTASE HOMOLOG [H.sapiens]
0553	-	AA028904	Hs.33215	ESTs
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18063	-	AA180054	Hs.73677	ESTS
39820	-	H24085	Hs.25443	ESTs
7374	-	AA093378	Hs.101810	ESTs
13109	-	AA435838	Hs.7985	ESTs
19378		H19673	Hs.31670	ESTs
24325	-	W84733	Hs.3978	ESTs
22318	-	R60224	Hs.7065	ESTs
24249	-	W73069	Hs.12600	ESTs
16514	-	AA027946	Hs.44608	ESTs
21421	- .	R21741	Hs.23258	EST
8397	-	AA426178	Hs.71725	ESTs Weakly similar to !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!! [H.sapiens]
8412	-	AA428090	Hs.26102	ESTs
10072	-	R14782	Hs.31931	ESTs
10349	-	AA001908	Hs.18160	ESTs
14492	-	AA609635	Hs.27497	ESTs
14930	-	T92512		EST - T92512
15861	-	X81001	Hs.141503	Small inducible cytokine A5 (RANTES)
16706	-	AA043800	Hs.62645	ESTs
16744	-	AA045643	Hs.62866	EST
16950	-	AA062980	Hs.66960	ESTs
17836	-	AA155779	Hs.29759	ESTs Weakly similar to !!!! ALU SUBFAMILY SX WARNING ENTRY !!!! [H.sapiens]
18834	-	F10207	Hs.13269	ESTs
19178	-	H10992	Hs.100910	EST
19767	-	H54720	Hs.36941	ESTs
21341	-	R14959	Hs.21871	EST Moderately similar to ninein (M.musculus)
21466		R24518	Hs.23674	EST
21602	-	R36624	Hs.135137	ESTs
21748	-	R40697	Hs.21290	EST
21850	-	R43089		EST - RC_R43069
21891	-	R43590	Hs.13290	ESTs
21937	-	R44508	Hs.22653	ESTs
22006	-	R46244	Hs.23110	ESTs
22054	-	R49116	Hs.25067	EST
22222	-	R55042	Hs.106645	ESTs
22232	-	R59385	Hs.141053	Homo sapiens bestrophin (VMD2) mRNA alternatively spliced product complete cds
22383	-	R63463	Hs.23282	ESTs
22446	-	R67259	Hs.25968	EST
23103	-	T23939	Hs.7344	ESTs
23872	-	T94562	Hs.17338	EST
24291	-	W80642	Hs.16951	ESTs

```
ESTs Moderately similar to !!!! ALU SUBFAMILY SC WARNING ENTRY !!!! [H.sapiens]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              EST
ESTs
ESTs Weakly similar to IIII ALU SUBFAMILY SQ WARNING ENTRY IIII [H.saplens]
EST
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Homo saplens short form transcription factor C-MAF (c-maf) mRNA complete cds
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ESTs Highly similar to ALANINE AMINOTRANSFERASE [Homo sapiens]
ESTs Weakly similar to mTERF [H.sapiens]
ESTs
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Spectrin beta non-erythrocytic 1
ESTs Weakly similar to LIS-1 proteln [H.sapiens]
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Homo sapiens BAC clone RG118D07 from 7q31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               H.sapiens mRNA for Pirin isolate 1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 EST - T35529
                                                                                                                                                                                                                                                                                   AFFX-Dapx-3
                                                                                                                       EST
ESTs
ESTs
ESTs
                                                                                                                                                                                                                                                                                                                     Hs. 97769
Hs. 104778
Hs. 9853
Hs. 9873
Hs. 12482
Hs. 102302
Hs. 112705
Hs. 112705
Hs. 112705
Hs. 102624
Hs. 10
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Hs. 104287
Hs. 40342
Hs. 51262
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Hs.132872
Hs.38842
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Hs.107164
Hs.107725
Hs.25985
Hs.31235
Hs.6624
   Hs.21403
Hs.90695
Hs.91440
Hs.90930
Hs.65749
Hs.65749
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Hs.6995
239086
A4435835
R06424
R44210
T79942
Z39430
Z39668
AFX-
A4412498
AA412498
AA422498
AA4231469
AA4231469
AA48840
AA48840
AA48840
AA609606
H42037
H91660
N47952
T03170
AA46587
AA465834
AA48719
WZ6496
R53024
AA427510
Z38153
H91255
   24640
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ESTs	ESTS	ESTs	ESTs	ESTs	ESTs	ESTs Highly similar to COATOMER ZETA SUBUNIT [Bos taunis]	ESTs	EST	Human done 23932 mRNA sequence	ESTs Weakly similar to BENOMYL/METHOTREXATE RESISTANCE PROTEIN [Candida albicans]	Homo sapiens mRNA for KIAA0573 protein partial cds	EST RC_R52088	Homo sapiens FGF-1 intracellular binding protein (FIBP) mRNA complete cds	ESTs	ESTs	EST - RC_AA084412	ESTs	ESTs	Human mRNA for uKATP-1 complete cds	EST - RC_W73946	EST - RC_AA421144	ESTs	ESTs	ESTs	ESTs ,	ESTs	EST - RC_H52379	ESTs	ESTs	Homo sapiens p38beta2 MAP kinase mRNA complete cds	ESTs .	EST - RC_R72597	ESTs Weakly similar to unknown protein [H.sapiens]	ESTs	ESTs Weakly similar to weak similarity to IHSP90 [C.elegans]	ESTs	ESTs	EST	ESTs Highly similar to 40S RIBOSOMAL PROTEIN S27A [Homo saplens Cavia porcellus]
Hs.36291	Hs.19865	Hs.124800	Hs.29126	Hs.12292	Hs.8961	Hs.37482	Hs.17265	Hs.17117	Hs.86921	Hs.10432	Hs.80844		Hs.7768	Hs.26590	Hs.61199		Hs.34183	Hs.116415	Hs.102308			Hs.79592	Hs.104441	Hs.16917	Hs.34274	Hs.24642		Hs.9899	Hs.72146	Hs.57732	Hs.87068		Hs.124570	Hs.5244	Hs.23294	Hs.72733	Hs.72499	Hs.112893	Hs.25996
H48825	AA401809	W87280	R77631	F10542	W26105	W61319	193870	T93078	U79257	L44334	R74235	R52088	C21105	R58922	AA024494	AA084412	R88711	AA609189	D50312	W73946	AA421144	H70121	AA281765	N67553	AA149826	AA291269	H52379	AA037199	AA156596	AA302831	AA232648	R72597	H12448	F09988	AA464689	AA180352	AA164750	AA620736	R71892
-	-	-	-	-	-	-	-	-	-	-		-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
19689	12450	24368	22565	18872	15358	24186	23863	23846	15143	9711	22544	41506	39345	22272	16434	17255	22692	38830	9179	42547	36195	29355	34608	20779	11081	12151	39935	7157	17858	34885	18445	22524	19224	18803	13810	18070	17937	39115	22517

2			1770000	
14251	-	AA497049	Hs.32501	ESTs
23923	-	T96407	Hs.17812	ESTs
21177	-	R06569	Hs.19721	ESTs
25705	-	AA131921	Hs.71030	ESTs
19805	-	H57725	Hs. 124031	ESTs
19668	-	H47656	Hs.33991	EST
36693	-	AA432389	Hs.98702	ESTs
14036	-	AA482107	Hs.5473	ESTs
20859	-	N69825	Hs.16762	ESTs
23849	-	T93113		EST - RC_T93113
18265	-	AA207122	Hs.86316	ESTs
35275	_	AA398530	Hs.97363	ESTs
10262	-	R77869	Hs.110493	ESTs
21757	-	R40789	Hs.21299	ESTs
21541	-	R31607	Hs.24420	ESTs
16873	-	AA056258	Hs.63264	EST
19646	-	H46006	Hs.31677	ESTs
23719	-	T89160	Hs.138506	ESTs
19608	-	H41581	Hs.31582	ESTs Moderately similar to HYPOTHETICAL 59.1 KD PROTEIN ZK637.1 IN CHROMOSOME III (Caenorhabditis elegans
17382	-	AA102731	Hs.109072	ESTs
16864	-	AA055971	Hs.63238	ESTs
10897	-	AA111881	Hs.9192	ESTs
17028	-	AA070178		EST - RC_AA070178
19220	-	H12318	Hs.24324	ESTs
9726	-	L44574	Hs.125235	ESTs
24570	0	Z38681	Hs.27150	ESTs
22167	ဂ	R53021	Hs.25873	ESTs
42537	9	W73417	Hs.103183	EST
18806	0	F10005	Hs.12599	ESTs
16585	0	AA033948	Hs.4236	ESTs
17309	0	AA086232	Hs.68717	EST Moderately similar to mariner transposase (H. sapiens)
22813	0	R97419	Hs.35718	ESTs
16429	0	AA022953	Hs.61172	EST
22013	0	R46526	Hs.25377	EST
8439	0	AA431277	Hs.32419	ESTs
22934	0	T10042	Hs.4205	ESTs
13063	0	AA432386	Hs.1191	Human mRNA for KIAA0073 gene partial cds
10122	0	R31745		EST - R31745
18195	0	AA195263	Hs.86001	EST

ESTs	ESTs	EST ·	ALPHA-2-MACROGLOBULIN PRECURSOR	ESTs	EST - BC AA079306
Hs.57836 ESTs	Hs.144270	Hs.86902	Hs.74561	Hs.95044	
W72557	AA063378	AA223929	AA219304	AA011210	AAATORAG
0	0	0	0	0	•

33249 16966 18363 34154 16222 17174

Primary Key	fold upragulated of Tumor over normal	ACCESSION	Unigeno Descriptor	ORF structural info
16074	colon >10	AA001507	ESTs	other
25047	>10	AA011031	ESTs	OCHEF
25062	>10	AA017257	EST:	other
16490	>10	AA026418	EST:	other
25179	>10	AA031268	H.sepiens mRNA for kinesin-2	other
25215	>10	AA035540	APOLIPOPROTEIN AI REGULATORY PROTEIN-1	other
25282	>10	AAD44825	EST:	other
16810	>10	AA053636	EST:	other
16835	>10	AA054438	EST•	SS.
10747	>10	AA055841	EST:	other
10748	>10	AA055892	EST:	other
16993	>10	AA065300	Human mRNA for KIAA0035 gene partial cds	other
17051	×10	AA070801	EST ₆	тм
10840	>10	AA084104	ESTs ·	other
7296	>10	AA085661	ESTs	other
7325 ·	»10	AA090842	ESTs Weakly similar to HYPOTHETICAL 35.7 KD PROTEIN C2F7 02C IN CHROMOSOME I [Schizosaccharomyces pombe]	other .
17419	>10	AA113349	EST	?
17541	>10	AA127459	ESTs	other
17559	>10	AA128407	ESTs	other
25669	>10	AA128978	ESTs	?
17600	>10	AA130596	EST:	other `
10992	>10	AA132523	Homo sapiena BAC clone RG119C02 from 7p15	other
17654	>10	AA133250	EST.	тм
17734	>10	AA137246	ESTs	ather
25801	>10	AA148530	EST -RC_AA148530	. TM
25806	>10	AA149007	EST	7
11121	>10 >10	AA156359 AA164289	Human TAR DNA-binding protein-43 mRNA complete cds ESTs	other
25925	>10	AA164494	ESTs Weakly similar to CYCLIC-AMP-DEPENDENT	other
25925	>10	AA165355	TRANSCRIPTION FACTOR ATF-6 [H.sapiens] Human clone lots unknown protein mRNA complete cds	7
17987	>10	AA169379	EST#	other
18008	>10	AA171895	Homo sapiens clone 24749 and 24750 mRNA sequences	TM
33953 33980	>10 >10	AA173290 AA180223	Human HOXA1 mRNA long transcript and alternatively speced forms complete cds CAMP-DEPENDENT PROTEIN KINASE TYPE II-	other
34013	>10	AA190888	ALPHA REGULATORY CHAIN EST - RC_AA190888	other
34014	>10	AA191348	ESTs .	other
34015	>10	AA191353	ESTs	TM
34070	>10	AA196549	ESTs	other
18260	>10	AA206801	ESTs	7
34105	>10	AA207123	ESTs	\$\$.
34107	>10	AA209469	EST:	other
18330	>10	AA216722	Human mRNA for apolipoprotein E receptor 2 complete	TM
10362	>10	AA223912	cds Ribonuclease L (25'-oligoisoxdenytate synthetase- dependent) inhibitor	other
18392	>10	AA227751	ESTs	\$ 5,
34158	>10 °	AA228030	EST:	TM
34197	>10	AA232315	Home sapiens close 23797 and 23917 mRNA partial cds	other
25948	>10	AA234365	Homo sapiens survival of motor neuron protein interacting protein 1 (SIP1) mRNA complete cds	?
25951	>10	AA234556	EST	,

FIGURE 8
1 of 37

11561	>10	AA236533		Evr1	other
26059	>10	AA236685		EST:	other
26100	>10	AA242835		Human mRNA for KIAA0391 gene complete cds	other
11603	>10	AA243052		ESTs Highly similar to GONADOTROPIN-RELEASING HORMONE RECEPTOR [Rattus norvegicus]	other
7785	>10	AA243375		EST - AA243375	other
34372	>10	AA251973		EST ₆	ş
26240	>10	AA252282		Human mRNA for KIAA0152 gene complete cds	TM
34382	>10	AA252512		ESTs	other
34391	>10	AA252703		EST - RC_AA252703	other
26274	>10	AA253011		EST:	,
34400	>10	AA253400		ESTs	other
26291	>10	AA253422		ESTs	?
26355	>10	AA256379		ESTs .	other
11813	>10	AA258158		ESTs	other
34518	>10	AA278721		EST:	,
26545	>10	AA278979		ESTs ·	other
26574	>10	AA279504	٠,	EST:	other
34554	>10	AA280016		DNA polymerase gamma	other
34561	>10	AA260298		ESTs .	TM
26628	>10	AA280641	*	ESTs Highly similar to 60S RIBOSOMAL PROTEIN L39	TM
11969	>10	AA280670		(Rattus norvegicus) ESTs	SS.
34575	. >10	AAZ80738		ESTs	TM
26677	>10	AA281636		ESTs	7
26700	>10	AA282197		EST	,
34672	>10	AA284372		EST:	other
34692	>10	AA285079		ESTs	piher
12143	>10	AA290991		ESTs	other
8092	>10	AA316272		ESTs .	TM
34904	>10	AA321746		EST	other
8111	>10	AA323787		ESTs	other
8125	>10	AA330771		Human protein-tyrosine phosphatase (HU-PP-1) mRNA	TM
26916	>10	AA331393		partial sequence ESTs	other
26926	- >10	AA342402		ESTs .	other
26935	、>10	AA347193		ESTs Weakly similar to NADH-UBIQUINONE	TM
35038	>10	AA350541		OXIDOREDUCTASE CHAIN 4 [Caeromabditis elegans] ESTs Moderately similar to URACIL-DNA	TM
35049	>10	AA350857		GLYCOSYLASE 1 PRECURSOR [H.sapiens] ESTs	other
35106	>10	AA371561		EST Weakly similar to putative p150 (H.sapiens)	7
35197	>10	AA398120		ESTS	other
35277	>10	AA398536		EST6	other
35309	>10	AA398660		EST	other
35322	>10	AA398710		H. sapiens RNA for CLCN3	TM
27037	>10	AA400198		ESTs	TM
35485	>10	AA400527		EST:	7
27046	>10	AA400670		Home sapiens mRNA for KIAA0582 protein partial cds	other
35500	>10	AA400715		ESTs	other
12480	>10	AA403116		Homo sapiena U-snRNP-associated cyclophilin (USA-	other
35693	>10	AA405485		CyP) mRNA complete cds ESTs Weakly similar to similar to a complex testis- specific protein (C. elegans)	other
35697	≻10	AA405512		ESTs	other
35766	>10	AA406169		Homo saprens KIAA0431 mRNA partial cds	other
35769	>10	AA406206		ESTs	other
35798	>10	AA410231		EST:	other

FIGURE 8 (cont.)
2 of 37

35801	>10	AA410291	EST6	Other
35803	>10	AA410295	EST:	other
35822 -	>10	AA411144	ESTs	TM
35874	>10	AA412024	EST	7
35958	>10	AA412550	EST•	attrer
36052	>10	AA417027	EST	TM
36258	>10	AA423962	ESTs Weakly similar to IIII ALU SUBFAMILY J	other
36288	>10	AA424502	WARNING ENTRY IIII (H.szpiens) ESTs	olher
36307	>10·	AA424803	EST	2
36371	>10	AA426017	ESTs	TM
36395	>10	AA426353	ESTs	other
36405	>10	AA426406	Homo sapiena mRNA for KIAA0530 protein partial cds	other
36506	>10	AA429610	EST ₈	other
36571	×10 °	AA430726 .	EST - RC_AA430726	SS.
36695	>10	AA433910	ESTs	other
36739	>10	AA435610	EST - RC_AA435610	7
36753	×10	AA435686	EST:	other
36845	>10	AA436198	EST:	7
13136	>10	AA436560	EST:	SS.TM
13143	>10	AA435619	ESTs	SS,
36958	>10	AA442060	ESTs	other
36962	> 10	AA442062	EST:	7
36961	>10	AA442845	EST	7
13237	>10	AA443971	ESTs Weakly similar to !!!! ALU SUBFAMILY J	7
13242	>10	AA445994	WARNING ENTRY IIII [H.sapiens] ESTs	other
37057	>10	AA445131	ESTs	other
37068	>10	AA445312	ESTs Weakly similar to 199 ALU CLASS C WARNING	other
37074	>10	AA446344	ENTRY IIII (H. sapiens)	SS.
37084	>10	AA446488	Homo sapiens Ran binding protein 2 (RenBP2alpha)	2
			mRNA partial cds	,
37135	>10	AA447540	EST	other
37159	>10	AA447714	EST - RC_AA447714	7
37168	>10	AA447772		: other
37246	>10	AA449311	Homo sapiens mitotic checkpoint protein kinase (BUB1) mRNA complete ods	•
37310	>10	AA451707	ESTs	other
37453	>10	AA454610	ESTs	7
37456	>10	AA454632	ESTs	TM
27787	>10	AAA54660	ESTs	7
37492	>10	AA455248	EST -'RC_AA455248	other
37546	>10	AA456641	ESTs .	TM
37601	>10	AA458864	ESTs	other SS.TM
37611	>10	AA458996	Human signaling lymphocytic activation molecule (SLAM) mRNA complete cds	
37615	>10	AA459101	Human sersie/threonine kinase mRNA partial cds	other
37653	>10	AA460017	ESTs	other
37677	>10	AA460530	Home sapiens orphan G protein-coupled receptor HG38 mRNA complete eds	other
37777	>10	AA464860	Homo sapierer Jak2 kinase mRNA complete cds	other
8648	>10 ·	AA465016	Homo sapiens serine protease-like protease (nes1) mRNA complete cds	7
37816	>10	AA469954	EST .	,
37829	>10	AA470084	ESTs	other
28015	>10	AA477421	ESTs ·	other
37978	>10	AA479294	EST - RC_AA479294	other
37979	>10	AA479295	ESTs Highly similar to RING CANAL PROTEIN [Drosophile melanogester]	other

FIGURE 8 (cont.) 3, of 37

37983	>10	AA479346	H sapiens mRNA for SYT	otiver
14054	>10	AA485223	EST:	TM
38121	>10	AA485724	EST - RC_AA485724	other
25122	~10	AA485928	ESTs Weakly similar to LOK (M.musculus)	other
38167	>10	AA487207	EST - RC_AA487207	other
38172	>10	AA487424	EST - RC_AA487424	other
38179	>10	AA487492	Homo sapiens clone 23592 mRNA sequence	other
38182	>10	AA487501	ESTs	other
38194	>10	AA487969	ESTs	other
28141	>10	AA488432	EST:	?
38211	>10	AA488687	ESTs .	other
30235	>10	AA489030	ESTs	other
38280	>10	AA489791	EST - RC_AA489791	other
38316	>10	AA490500	Homo sepiens ribonucleese P protein subunit p20	other
38330	>10	AA490882	(RPP20) mRNA complete cds ESTs	other
38456	>10 .	AA504343		SS.
38460	>10	AA504462	ETT:	other
38553	>10	AA521471	EST:	other
38580	>10	AA598545	ESTs	?
38590	>13	AA598648	Human mRNA for transcriptional activator hSNF2b	other
			complete cds	2
39601	>10	AA598738	ESTs	•
	>10	AA599639	ESTs .	other
38825	>10	AA609177	ESTs .	TM 2
38838	>10	AA609215	EST - RC_AA609215	TM
38867	>10	AA609318	Human ctil-b mRNA complete cds	7
38871	>10	AA609333	EST	other
38970	>10	AA609749	ESTs	ouner ?
38964	>10	AA609839	ESTs Moderately similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII [H.sapiens]	•
39045	>10	AA610077	EST:	other
39062	>10	AA620333	EST	?
39080	>10	AA520552	EST - RC_AA620552_r	7
39110	>10	AA620709	ESTs Weakly similar to HYPOTHETICAL 90.6 KD PROTEIN T05H10.7 IN CHROMOSOME II (C.elegans)	other
39176	>10	AA621091	EST9	other
39218	>10	AA621330	EST#	other
39221	>10	AA621346	Homo sapiens protein phosphatase with EF-hands-1 (PPEF-1) mRNA complete cos	other
39232	>10	AA621409	EST:	olhar
21	>10	AB000905	H.sapiens histone H4 gene	7
89 63	>10	AFFX- HUMTFRRM115 07	AFFX-HUMTFRR/M11507_M	7
33890	>10	AFFX- HUMTFRR/M115	AFFX-HUMTFRRAM11507_5	7
39302	>10	C14944	ESTs	other
39329	>10	C20797	EST	7
26544	>10	D12163	ESTs	other
218	>10	D13540	PROTEIN-TYROSINE PHOSPHATASE 2C	other
236	>10	D13645	Human mRNA for KIAA0020 gene complete cds	other
9127	>10	D30037	PHOSPHATIDYLINOSITOL	other
459	>10	D38293	Human mRNA for clatterin-like protein complete cds	TM
39405	>10	D50975	ESTs	other
39433	>10	DS2037	Human thymidine kinase 2 (TK2) mRNA complete cds	other
39436	>10	D52692	Human Ca2+-dependent activator protein for secretion	TM
14706 .	>10	D59388	mRNA complete cds	,

FIGURE 8 (cont.)
4 of 37

WO 00/55633 PCT/US00/07044

39488	>10	D60831	ESTs .	other
39504	>10	080632	ESTs .	other
765	>10	D86096	Prostaglandin E receptor 3 (subtype EP3) (alternative products)	7
787	>10	D86969	Human mRNA for KIAA0215 gene complete cds	other
789	>10	D86971	Human mRNA for KIAA0217 gene partial cds	other
39529	>10	F02202	ESTs	7
39535	>10	F02450	ESTs Moderately semilar to unknown protein (H. sapiens)	TM
18678	>10	F04022 .	EST#	other
18718	>10	F04915	ESTs.	other
18762	>10	F09458	EST#	other
18782	>10	F09739	EST•	other
29080	>10	F13655	ESTs Moderately similar to fill ALU SUBFAMILY SB1 WARNING ENTRY fill (H.sapiens)	other
19001	>10	H02890	ESTs	other
19164	>10	н10395	TEST	?
39725	>10	H11323	EST:	other
19203	>10	H11593	EST:	other
19328	≯10	H17808	EST:	other
19387	>10	H20128	ESTA	7
39787	>10	H20131	ESTs	\$5.
19389	>10	H20165	EST	?
39832	<u>>10</u>	H26279	EST - RC_H26279	other
19591	>10	H40688	EST:	other
29229	>10	H48459	Human mRNA for KIAA0186 gene complete cds	other
19727	>10	H52702	EST&	?
19787	>10	H56679	EST\$	other
39995	>10	H62474	EST	SS.TM
29331	>10	H68116	ESTE	TM
29344	>10	H68839	EST	7
40064	>10	H72283	Human mRNA for KIAA0265 gene partial ods	other
40083	>10	H73466	MITOCHONDRIAL 60S RIBOSOMAL PROTEIN L3	other
19949	>10	H78263	ESTs	TM
40204	>10	н88296	EST - RC_H88296	other
29523	>10	H88353	ESTs Weakly similar to line-1 protein ORF2 [H.sapiens]	7
29551	>10	H90134	ESTs	7
29645	>10 .	H95840	EST:	7
20057	>10	H98079	EST.	ather
976	>10	HG2036-HT2090		7
1158	>10		EST - HG3344-HT3521	7
1210	>10	HG37-HT37	EST - HG37-HT37	7
1346	>10	HG4718-HT5158	•	7
1349	>10		EST - HG4747-HT5195	,
1445	>10	J03027	MHC class I protein HLA-G	?
1570	>10	K01383	EST - K01383	•
1664	>10	L07541	Replication factor C (activator 1) 3 (38kD)	other TM
1852	>10	L17328	Human FEZZ mRNA partial cds	1M ?
1856	>10	L18920	MELANOMA-ASSOCIATED ANTIGEN 2	? other
1863	>10	L19161	TRANSLATIONAL INITIATION FACTOR 2 GAMMA SUBUNIT	
2070	>10	L37378	Homo sapiens guenytyt cyclase (RetGC-2) mRNA complete cds	SS,TM
2123	>10	L40396	Homo sepiens (clone s22:71) mRNA fragment	other
2144	>10	L41349	Phospholipase C beta 4	\$ \$.
9723	>10	L44542	EST#	other
2188	≯10	L47276	EST - L47276	other

FIGURE 8 (cont.)
5 of 37

2343	>10	M15353	Eukaryotic translation instation factor 4E	other
2627	>10	M29810	Glycophorn E	TM .
2857	>10	M58597	Fucosytransferase 4 (alpha (13) fucosytransferase myeloid-specific)	TM
3021	>10	M68941	Protein tyroses phosphatase non-receptor type 4	othe:
3163	>10	M84424	Cathepsin E	7
3196	>10	M86917	Oxysterol binding protein	other
20088	>10	N20054	ESTs Weakly similar to putative p150 [H.sapiens]	7
29795	>10	N20641	EST& Highly similar to HYPOTHETICAL MYELOID	other
40427	>10	N21147	CELL LINE PROTEIN 3 [Hamo sapiens] ESTs	other
40444	>10	N22140	ESTs Highly similar to TUBULIN GAMMA CHAIN	other
29893	>10	N23003	[Euplotes octocarinatus] ESTs	TM
40498	>10	N26086	Homo sapiens citrate synthase mRNA complete cds	SS.
	>10	N33024	ESTs	SS,
40559	>10	N33264	EST	7
30190				
30207	>10	N33920	H.sspiens mRNA for diubiquitin	other
20304	>10	N34686	Home sepiens clone 23915 mRNA sequence	
20307	>10	N34830	ESTs	other
30265	>10	N35115	ESTs	other
40594	>10	N35388	EST:	other
40604	>10	N36893	Homo sapiens KIAAD428 mRNA complete ods	other
40631	>10	N45124	ESTa	other
40660	>10	N43104	NUCLEAR FACTOR RIP140	other
30510	>10	N50138	EST	?
30617	>10	N50646	EST:	other
30631	>10	N50807	EST	7
30790	>10	N52935	EST	7
20554	>10	N55443	ESTs	TM
40760	>10	N57927	ESTs Weakly similar to ELL [M.muscutus]	other
30938	>10	N58561	Cathepsin 8	other
20614	>10	N59230	ESTs	55 .
20657	>10	N52889	EST«	other
31136	>10	N63512	ESTs Weakly similar to M01F1.4 [C.elogans]	TM
40827	>10	N54051	Homo sapiens Werner syndrome gene complete cda	other
31310	>10	N66831	EST	7 .
40876	>10	N67607	Human Rho-exsociated colled-coil containing protein	other
20791	>10	N68057	kinase p160ROCK mRNA complete cds Home saplens telement repeat binding factor (TRF1)	7
40905	>10	N58738	mRNA complete cds ESTs	other
40905	>10	N59114	H.sapiens mRNA for orphan nuclear hormone receptor	other
40913	>10	N59218	· ESTs	other
31484	>10	N69466	EST:	CURE
31619	>10	N73449	ESTs	other
41005	>10	N79516	EST:	TM
31818	>10	N89774	Homo sapiens mRNA for KIAA0292 gene partial cds	giner
31872 41040	>10 ·	N91109 N91948	EST EST#	other
41040 31944	· -		ESTS	•
	>10	N93183		other
41065	>10	N93618	, EST4	other
32034	>10	N98926	ESTs Moderately similar to DMR-N9 PROTEIN [H,sepiens]	other
41107	>10	R01634	EST:	other
41163	>10	R08176	EST ₆	other
21238	>10	R08564	Plasminogen-lika protein	other

FIGURE 8 (cont.)
6 of 37

21240	>10	R08613	ESTs	other
21412	>10	R20670	ESTs	other
21519	>10	R27975	EST - RC_R27975 .	other
41361	>10	R42278	H sapiens mRNA for TRES	7
32189	>10	R43183	ESTs	TM
32195	>10	R43471	ESTs Wealdy similar to ORF YOR258w [S.cerevisize]	other
21902	>10	R43822	EST	7
21946	>10	R44707	EST#	MT
22072	>10	R49406	EST»	7
32240	>10	R50976	Ribonuclease L (2'5'-oligorsoadenylate synthetase-	other
32258	>10	R55823	dependent) inhibitor ESTs	other
22258	>10	R56432	EST.	other
22282	>10	R58197	ESTs	other
32277	>10	R61493	Human mRNA for rod photoreceptor protein complete	other
22372	>10	R62831	cots EST	,
22400	>10	R64109	ESTs	other
41593	>10	R64129	ESTs	other
10233	>10	R71427	ESTs Highly similar to PHENYLALANYL-TRNA	other
10233	0		SYNTHETASE ALPHA CHAIN CYTOPLASMIC (Saccharomyces cerevisize)	
41654	>10	R76437	THROMBOXANE-A SYNTHASE	TM
22557	>10	R76722	EST#	other
22576	>10	R79111	ESTs	other
22593	>10	R79777	EST	7
41678	>10	R80675	EST	7
41719	>10	R89260	EST - RC_R89260	other
22793	>10	R96208	ESTs	other
41752	≥10	R97063	EST\$	other
3375	>10	\$50223	нкя-т1	other
3406	>10	S66896	SQUAMOUS CELL CARCINOMA ANTIGEN 1	other
3522	>10	S80267	Spleen tyrosine kinase	other
41793	>10	T03887	ESTs	?
23198	>10	T40530	ESTs Weakly similar to B0035.14 [C.elegans]	other
23360	>10	T58531	ESTs	other
32635	>10	T61116	ESTs	other
42177	>10	T79786	EST	7
23623	>10	T84047	ESTs .	7
23662	>10	T86674	ESTs .	other
42242	>10	T69579	Homo sapiens E2F-related transcription factor (DP-1) mRNA complete cds	other
23759	>10	T90313	ESTs	other
23832	>10	T92018	EST:	other
32740	>10	T92950	EST:	other
42290	>10	T95105	ESTs	7
3598	>10	U01157	Glucagon-like peptide-1 receptor	SS.TM
3659	>10	U04313	Protease inhibitor 5 (maspin)	other
3799	· >10	U10690	Human MAGE-5s antigan (MAGE5e) gene complete	7
3870	>10	U14518	cots Centromere protein A (17kD)	other
3913	>10	U16261	Human MDA-7 (mda-7) mRNA complete cds	5\$.
4029	>10	U21090	Human DNA polymerase delta smell subunit mRNA	other
4157	>10	U28811	complete cds Human cysteine-rich fibroblest growth factor receptor	other
****	>10	U30246	(CFR-1) mRNA complete cds	TM
4178			Human bumatanide-sensitive Na-K-CF cotransporter (NKCC1) mRNA complete cds	
15006	>10	U30246	Human bumetanide-sensitive Na-K-Cl cotransporter (NKCC1) mRNA complete cds	TM

FIGURE 8 (cont.)
7 of 37

4193	>10	U31116	Human beta-sarcoglycan A3b mRNA complete cds	TM
4306	>10	U36798	Homo sepiens platelet cGI-PDE mRNA complete cos	TM
4362	>10	U39817	Bloom syndrome	other
4386	>10	U40622	DNA repair protein XRCC4	other
4388	>10	U40714	Human tyrosyl-IRNA synthetase mRNA complete cos	other
4455	>10	U43944	MALATE OXIDOREDUCTASE	other
4477	>10	U45880	Human IAP-like protein ILP mRNA complete cds	other
4680	>10	U55766	Human Rev Interacting protein Rip-1 mRNA complete	TM
4702	>10	U57341	EST - U57341	other
4713	>10	U57721 .	Human L-kynurenine hydrotase mRNA complete cds	other
4787	>10	U61145	Human enhancer of zeste homolog 2 (EZH2) mRNA	otner
4862	>10	U65437	complete ods Human homeodomein-containing protein (HANF) mRNA	>
4945	>10	U69108	complete cds Home sapiens mRNA for TRAFS complete cds	other
				pliner
4975	>10	U71088	Human MEK5 mRNA complete cds	other
4994	>10	U72514	Human C2/ mRNA complete cds	other
5002	>10	U72761	Human karyopharin bata 3 mRNA complete ods	TM
5021	>10	U73524	Human putative ATP/GTP-binding protein (HEAB) mRNA complete cds	
5149	>10	U79716	Human reciin (RELN) mRNA complete cds	SS.
5214	>10	U83303	H.sapiens mRNA for granulocyte chemotactic protein	?
5243	>10	U85946	Human brain secretory protein hSec10p (HSEC10) mRNA complete cds	other
32789	>10	W02779	ESTs Moderately similar to kinezin-73 [D.melanogaster]	other
42354	>10	W19346	ESTs .	ather
42390	>10	W40150	Homo sapiens chromosome-essociated polypeptide	other
33006	>10	W46286	(HCAP) mRNA complete cds ESTs Weakly similar to ZK1058.5 [C.elegans]	TM
33020	>10	W46891	ESTs Weakly similar to polypeptide N-	other
33109	>10	W59961	acetytgalactosaminyliraristerase (H.sapiens) Human mRNA for KIAA0389 pena complete cds	olher
24197	>10	W67277	ESTs	other
24215	>10	W69425	ESTs	other
33301	>10	W73883	FSTA	ather
33343	>10	W79834	ESTs Weekly similar to rhotekin [M musculus]	other
33377	>10	W81219	ESTs Weakly similar to F4686.7 [C.elegans]	other
42602	>10	W86423	ESTS (VERKIY SATINGS TO PAGGO. 7 (C. GEFGATS)	TM
33556	>10	W90705	Murine loukemis viral (brsi-1) oncogene homolog	other
33616	>10	W93726	Protease inhibitor 5 (maspin)	other
33666	>10 >10	W95876	ESTs .	TM
5510	>10	X05360		2
5558	>10	X05360 X07876	Cell division cycle 2 G1 to S and G2 to M Wingless-type MMTV integration site 2 human homolog	SS.
5603	>10	X1/8/6 X14253	Taratocarcinoma-derived growth factor 1	TM
5619	>10	X14850	HISTONE HZA.X	SS
5623	>10	X14975	T-CELL SURFACE GLYCOPROTEIN CD1E	,
			PRECURSOR	
5692	>10	X17644	G1 to S phase transition 1	other
5789	>10	X54925	Matrix metalloproteinase 1 (interstitial collagenase)	other
5799	>10 -40	X55330	Aspartylglucosaminidase	\$8.
5802	>10	X55544	CYCLIC-AMP-DEPENDENT TRANSCRIPTION FACTOR ATF-1	•
5857	>10	X58377	Human mRNa for adipogenesis inhibitory factor	other
5960	>10	X63575	ATPase Ca++ transporting plasma membrane 2 (NOTE: redefinition of symbol)	TM
5963	>10	X63629	Cadherin 3 (P-cadherin)	SS.TM
5986	>10	X64810	Proprotein convertase subtilisin/kexin type 1	7
6041	>10	X67155	MITOTIC KINESIN-LIKE PROTEIN-1	ather
6095	>10	X69962	Fragile X mental retardation 1	other

FIGURE 8 (cont.) 8 of 37

6106	>10	X70683	SRY (sex determining region Y)-box 4	TM
6155	>10	X74331	DNA primese polypoptide 2A (S8kD)	19/30
6167	>10 ·	X74987	Ribonuciease L (2'5'-oligoisoadenylate synthetase- dependent) inhibitor	other
5168	>10	X76029	NEUROMEDIN U-25 PRECURSOR	TM
6315	>10	X81889	H.szpiens mRNA for p0071 protein	other
6382	>10	X85133	H.sapiens RBQ-1 mRNA	other
6384	>10	X85137	Human kinesin-like spindle protein HKSP (HKSP) mRNA complete ods	other
6438	>10	X89396	URACIL-DNA GLYCOSYLASE 1 PRECURSOR	?
6449	>10	X89986	H.sapiens mRNA for NBK apoptotic inducer protein	TM
6478	>10	X91648	H.sapiens mRNA for pur alpha extended 3'untranslated region EST - X91653	SS.TM
6479	>10	X91653	H.sapiens mRNA for UDP-GatNActpolypeptide N-	TM
6494 - 6713	>10 >10	X92689 Y08564	ecetylgalactoseminyl transferase EST - Y08564	7
	>10	Y12394	Homo sepiens importin-style homolog (SRP1gamma)	other
6790 24915	>10	YEL003w/	mRNA complete cds : EST - YEL003w/	?
42773	>10		EST - YEL019c/MMS21	7
24545	. >10	Z38462	EST:	other
33713	>10	Z39427	ESTs	other
33791	>10	Z40883	EST4	other
42766	>10	Z99394	ESTs Moderately similar to IIII ALU SUBFAMILY SP	other
21558	>10	R33112	WARNING ENTRY (III [H.sapiens] Human AF-6 mRNA complete cds	other
26718	>10	AA282576	EST:	7
40113	9.9955090946	H78003	ESTs	7
10801	9 9979448276	AA069285	ESTs Weskly similar to PROBABLE UBIQUITIN CARBOXYL-TERMINAL HYDROLASE R10E11.3	other
37491	9.9513600842	AA455239	(C.elogans) ESTS Highly similar to CHROMOSOME CONDENSATION PROTEIN DPY-27 (Csenorhabditis elogans)	other
23900	9.9272347693	T95789	ESTs	cther
254	9.9198395324	D14657	Human mRNA for KIAA0101 gene complete cds	other
6885	9.6970927914	Z29331	Ubiquitin-conjugating enzyme E2H (homologous to yeast UBC6)	other SS
29693	9.8850766398	H97819	ESTs	
26482	9.8765189024	AA262491	ESTs	other 2
23123	9.8699502035	T25306	EST	other
26525	9.8160399123	AA276392	ESTs	
13110	9.7643356605 9.7087597628	AA435840 AA299784	Homo sepiens mRNA for high mobility group protein HMG2e EST	other
39432	9,7034550083	O51691	Phosphoribosylglycinamide formyltransferase	1
31312	9.6513325388	N66845	phosphoribosylgiyolnamide synthetase phosphoribosylaminoimidazole synthetase ESTs Weakly similar to IIII ALU CLASS B WARNING	2
	9.6358446349	R01179	ENTRY (()) [H.sapions]	7
21112	***************************************			other
31572	9.6254820695	N71294 AA160759	EST:	7
17903 20747	9.6221229759 9.6094813734	N66842	ESTs	other
4676	9.589223908	U55206	Homo sepions human gamma-glutamyl hydrolase (hGH)	TM
34363	9.5627081023	AA251587	mRNA complete cds Home expiers mRNA for KIAA0530 protein partial cds	other
39094	9.540768988	AA620636	EST:	other
3888	9.5372000133	U15128	Human beta-12-N-eostylgtucosaminytransferase (I	7
39366	9.506250529	D12184	(MGAT2) gene complete cds ESTs	TM
7674	9.4458059039	AA203742	EST:	other
4192	9.4329744134	U31099	Human DP prostanoid receptor (PTGDR) mRNA partial	TM
4507	9 422674945	U47050	cds Human putative calcium influx channel (http3) mRNA complete cds	тм

FIGURE 8 (cont.)
9 of 37

35606	9,412026255	AA402227	ESTs Moderately similar to N-tropomodulin [Rinorvegous]	other
4970	9 3649551013	U70862	Human nuclear factor I-B2 (NFIB2) mRNA complete cds	7
19829	9.3432151573	H58813	EST	7
14837	9.2878584141	T40145	ESTe	TM
17336	9.2622148675	AA099585	ESTs	other
40541	9.2532836505	N30160	ESTs	other
29498	9 2487643833	H85434	EST	7
29943	9 1797074262	N24766	ESTs Moderately senitar to IIII ALU SUBFAMILY J WARNING ENTRY IIII (H.sepiens)	Тм
17997	9.1629681314	AA169633	EST	other
21320	9.1243463318	R11673	EST:	other
13883		AA476917	ESTs Weatdy similar to No definition kne found [C.elegans]	other
30539	9.0886887776	N49072	EST:	other
32778	9.0877919549	W02063	EST	?
26380	9.0809559378	AA257012	EST	7
15888	9.0595893607	X95632	Human Abl interactor 2 (Abi-2) mRNA complete cds	other
40812	9.0012874244	N63419	ESTs	other
903	8.9640387908	D90070	ATL-derived PMA-responsive (APR) peptide	other
22674	8.9515777733	R87160	EST:	TM
49807	6.9510132281	N62995	TRANSCRIPTION INITIATION FACTOR IIF BETA	other
15244	8 9195644974	W00904	SUBURIT EST:	TM
32296		R67075	Zinc finger protein X-linked	other
18269		AA209467	EST4	other
19682	8.8507628284	H47391	EST:	other
41607	6.833925517	R67868	CLEAVAGE SIGNAL-1 PROTEIN	other
2548	6.6299864699	M25897	Platislet factor 4	TM
7736	8.8279341243	AA232121	Human tyrosyl-IRNA synthetase mRNA complete cds	other
34490		AA262354	ESTs	other
38658	8 7669313482	AA599477	ESTs	other
7528	8 765157554	AA149543	ESTS	other
39939	8.7555031142	H53454	EST - RC_H53454	other
25111	8 7232692309	AA020787	ESTs	other
21655	8,716167279	R38239	EST	?
39663	8.665962852	H04756	ESTs Highly similar to NADH-USIQUINONE	other
1042	8.652112324	HG2510-HT2806	OXIDOREDUCTASE 817 SUBUNIT (Bos taurus) EST - HG2510-HT2606	7
32330	8.6361115426	R77776	EST&	other
25382	8.6239456487	AA059007	ESTs	other
27074	8,5900813076	AA401475	ESTs Weakly similar to C3681.3 [C.slogans]	55.
3955	8.5298909183	U18259	MHC class II transactivator	other
4959	8 52646827	U70322	Human transportin (TRN) mRNA complete cds	other
2315	8.5259185808	M14123	EST - M14123_xp11	7
37253	8.4895914632	AA449357	EST:	other
39624	8,471316877	F10836	ESTS	7
23213	8.4569920887	T40691	ESTs	7
2798	8.455596435	M54995	Connective tissue ectavation peptide III	TM
41154	8.4413390141	R07499	ESTs	,
32479	8.4093689549	T16282	WEET-LIKE PROTEIN KINASE	other
41251	8,3587565415	R28279	Human clone 23548 mRNA sequence	other
19081	8.3583603183	H05701	ESTs Weakly similar to RHOMBOTIN-1 [H.sapiens]	other
21098	8.3105927559	R00545	EST:	other
14723	8.3061679053	D59894	ESTs	other
37154	8.2994822341	AA447666	Human CENP-F kinetochore protein mRNA complete ads	other

FIGURE 8 (cont.)
10 of 37

8068	6 2835586361	AA313387	ESTa Highly similar to HYPOTHETICAL 84 7 KD PROTEIN ZK1088.1 IN CHROMOSOME III (Ceenorhabditis elegans)	other
7485	8.281679348	AA129547	EST:	other
16501	8.2517969834	AA026969	EST.	other
34527	8.2419163754	AA279091	EST#	other
6700	8 1948675662	Y07867	H.sapiens mRNA for Pirin isolate 1	other
2852	8.1928816537	M58450	Human 75-kD autoantigen (PM-Sc1) mRNA complete cots	other
11188	8 1862492468	AA172372	ESTs	TM
42293	8.183311064	T95333	ESTs Weakly similar to coded for by C. elegans cDNA	TM
5443	8 1763317544	X02530	yk110g8,3 (C.elegans) Interferon (gamma)-induced cell line protein 10 from	SS,
40937	8.1534810594	N70607	EST:	TM
23371	8,1499496068	T59505	EST - RC_T59505	7
26272	8.1339974519	AA252981	ESTs Weakly similar to K07C11.10 gene product	other
	8 1332403762	AA086201	(C.slegans)	other
17306	4,1104,104,104	AA233795	EST:	other
18497	8.1192326373			other
235	8.0944363901	D13644	Human mRNA for KIAA0019 gene complete cds	TM
24525	8.0860187097	Z38347	****	TM
7826	8.0750029554	AA248884	EST - AA248884	other
32142	8.0739258775	R38715	Homo sepiens clone 24540 mRNA sequence	••••
39067	8.0557768803	AA620405	ESTs	other TM
6235	8.0448957236	X78416	Casein elpha S1	
29517	6.0017588725	H88261	ESTs .	other
28570	7.9852455973	C21104	Homo sapiens STAT-Induced STAT inhibitor-2 mRNA complete cds	other
39344	7.9162087762	C21034	ESTs Moderately similar to initiation factor afF-2B gamma subunit [R.norvegicus]	other
18951	7.9002189759	H00580	EST:	other
18953	7.8709160227	H00815	ESTs	other
18376	7.8564099916	AA226925	EST.	other
19830	7.847878447	H58911	EST4	other
36023	7.840835828	AA416881	EST:	'pther
13347	7.8344414518	AA449238	ESTs	other
36614	7.8284591351	AA431466	ESTs	other
2192	7.6254072032	L48211	Homo Sapiens angiotensin (I receptor gene complete cds	7
33016	7.6006574068	W46577	H sepiens mRNA for ESM-1 protein	other
17215	7.7941954038	AA083044	EST:	other
34894	7.7659738105	AA311881	EST	7
40814	-7.695001222	N39257	EST:	other
36295	7.6834749899	AA424534	ESTs .	other
19564	7.6744302788	H38833	EST:	TM
16914	7.6686405336	AA058665	ESTs	58.
35967	7.6378079107	AA412694	Human splicing factor SRp55-2 (SRp55) mRNA	olher
21672	7.6364823402	R38635	complete cds ESTs	other
19918	7.6303275831	H69787	ESTs	7
10511	7,6297744492	AA024482	ESTs Highly similar to KERATIN TYPE I	other
17721	7.6057911016	AA136590	CYTOSKELETAL 14 (Homo sapiens) ESTs	?
17721 42302	7.6057911016	T96130	EST	ss.
		196130 AA243763	EST:	other
26134 18766	7.6000619383 7.5621799008	F09497	ESTs	other
34492	7.501590494	F09497 FA262439		other
270	7.501590494 7.4512152125	D14822	ATL-derived PMA-responsive (APR) peptide EST • D14822	other
35975	7,4177746986	AA412738	EST:	other
29842	7.4095809671	N21688	EST:	7

FIGURE 8 (cont.)
11 of 37

35389	7 3913043319	AA399555	EST:	other
19979	7.3868157166	H98477	ESTs	other
5793	7 3865864025	X54942	CDC28 protein lunase 2	other
19978	7 380969715	H87770	EST - RC_H87770	other
1260	7.3691089318	HG4126-HT4396	EST - HG4128-HT4396	7
31571	7 3676263454	N71250	ESTs	other
23765	7.3541191734	T90443	ESTs Weakly similar to KIAA0376 [H.samens]	?
35123	7.3397933455	AA380927	EST	7
38252	7.3341119467	AA489247	ESTs	other
36216	7.3282021037	AA488861	EST6	other
29418	7.2489407005	H77915	EST - RC_H77915	7
4834	7,1980951054	U63541	Human mRNA expressed in HC/HCC livers and MofT-4	other
42504	7.1913036522	W59803	proliferating cattle partial sequence ESTs	other
6111	7,158000198	X71125 ·	H, kapiens mRNA for glutemine cyclotransferase	MT
41773	7,154479618	T03024	ESTs Weakly similar to ribosomal protein L7 (H.sapiens)	other
9951	7.1363626365	N71513	ESTs	other
28109	7.0941968224	AA485212	ESTs	other
988	7.0783044659	HG2150-HT2230	EST - HG2160-HT2230	7
29848	7.0610668511	N22107	EST#	other
30626	7.0607950168	N50744	ESTs	other
22567	7.0225726353	R77771	ESTs	TM
9347	7.006323071	H03686	ESTs ·	TM
11696	7.0026773299	AA252894	ESTs ·	olher
40584	7.0010096333	N34870	EST	?
193	6.9767029188	D10923	PROBABLE G PROTEIN-COUPLED RECEPTOR HM74	TM
18305	6.9740536051	AA214048	Collegen type IV alpha 4	other
6078	6.9699682397	X69141	FARNESYL-DIPHOSPHATE FARNESYLTRANSFERASE	other
26741	6.902658703	AA283198 '	ESTs	other
35069	6.8992885685	AA358397	EST	7
23504	6.8977135983	T71042	ESTs	DUNET
299	6.8824513029	D16815	Home sepiens orphan nuclear hormone receptor 8073	other
40583	6.8689903023	N34855	mRNA 3' end ESTs	other
31428	6.8623762224	N68594	ESTs	other
6169	6.6606959727	X75091	SET PROTEIN	other
39524	6.6567355171	F01905	MALATE OXIDOREDUCTASE	other
34578	6.8430889439	AA280837	ESTs	other
38678	6.837527995	AA599920	Small inducible cytokine A5 (RANTES)	other
23936	6.8251471804	T96930	ESTs	other
9326	6.8181321394	D89377	Msh (Orasophila) homeo box homolog 2	ather
19158	6.6067351968	H11255	ESTs Highly similar to ACTIN-LIKE PROTEIN [Bos	TM
18185	6,7882148811	AA194983	taurus] Homo sapiens mRNA for osteoclastoganesis inhibitory fector (OCIF) complete cds	other
27028	6.757529124	AA399630	ESTs Weskly similar to KIAA0371 [H.sapiens]	other
41289	6.7519531681	R37265	EST	other
34511	6.7364448798	AA278298	EST - RC_AA278298	other
1566	8.7056207716	J05614	EST - J05614	7
25675	6.6692299748	AA129757	ESTs Highly similar to 60S RIBOSOMAL PROTEIN L22 [Rattus norvegicus]	ather
5814	6.6584342828	XS6088	CYTOCHROME P450 VII	SS.
13861	6.6238291507	AA470145	ESTs	other
29794	6 6026313352	N20598	ESTs .	other
39333	6.5902382643	C20910	Cyclin B1	other
3770	6.5835303599	U09609	Nuclear factor of kappa light polypeptide gane enhancer in 8-cetts 2 (p49/p100)	other

Nuclear factor of kappa kight polypepti in B-celts 2 (p49ip100) FIGURE 8 (cont.) 12 of 37

31831	6.5829933764	N89894	ESTs	7
33063	6.5808125026	W53000	Homo sapiens clone 24431 mRNA sequence	other
20326	6.5640084836	N35583	ESTs Weakly smiler to PROSABLE E5 PROTEIN	7
34384	6.5535703492	AA252537	[Human papitomavirus type 58] ESTs	other
25599	6.5490481991	AA114091	Human (clone 881) Br-cedherin mRNA complete cds	other
39749	6.5369363254	H14968	ESTs	other
42596	6.5200567072	V/65900	ESTs	7
39606	6.5119482185	F10243	ESTs Wealdy synilar to till ALU CLASS 8 WARNING	7.
14617	6.5105504748	C14983	ENTRY (III (H.capiens)	other
27831	6.45670814	AA456044	EST:	7
34896	6.4496517783	AA312551	EST	7
27360	6,4434305006	AA425356	ESTs	other
20126	6.4326610424	N22015	ESTs	тм
6663	6.4324809977	Y00291	RETINOIC ACID RECEPTOR BETA-2	TM
30692	6.4196636207	N51563	ESTs .	other
36472	6.4189542265	AA428633	EST	7
9578	6.3961768753	H87652	Homo sapiens bicaudai-D (BICD) mRNA complete cds	other
39670	6.3818496159	H05626	ESTs	other
22697	6,3652792447	R89218	ESTs	other
37308	6.3647804993	AA451694	EST	TM
16101	6.3517262802	AA002147	EST	7
20629	6.3486854401	N59798	ESTS	other
36100	6.3364146287	AA417740	EST#	7
15488	6.3252590241	W28097	Homo sapisns clone 23711 unknown mRNA partial cds	other
36667	6 3131273544	AA432136	EST:	other
30766	6.3115037924	N52627	EST - RC_N52627	7
32882	6.2745311453	W37683	ESTA	TM
18072	6.2675797205	AA180448	EST	7
16231	6.2652604863	AA199747	Human mRNA for KIAA0096 gene partial cds	other
38282	6.2514165678	AA489814	EST	7
28125	6.250317021	AA486073	ESTs	other
37464	6.2484456382	AA454747	ESTs	7
36618	6 1946328223	AA431478	ESTs	other
5082	6.1931116815	U78524	Human Gu binding protein mRNA partial cds	other
1441	6.1777287039	J02963	Integrin sipha 2b (plateist glycoprotein ilb of lib/lila	other
42105	6.14875944	T67710	complex entigen CD41B) ESTs	7
6061	6,1394863141	X68314	Giutathione peroxidase 2 gastrointestinat	SS.
32570	6.1156028796	T30222	ESTs Weakly similar to tetracycline transporter-like	тм
32504	6.1019612076	T17063	protein (M.muscutus) EST	,
23335	6.0977927504	T56804	EST	7
10967	6.0970991075	AA088458	ESTs Weakly similar to IIII ALU SUBFAMILY J	other
			WARNING ENTRY III! (H.sapiens)	. 7
30863	6.0911993489	N56923	EST .	TM
14528 29454	6.0859008453 6.0685955038	AA620295 H81306	EST .	7
6798	6.0539173278	Y13153	Homo segrens mRNA for kynucenine 3-monodxygenese	m.
21248	6.0539173278	R06871	ESTs	·m
21940	6.0499964138	R44538	EST4	,
29066	6.0455247653	F10927	Homo sepiens clone 23636 mRNA sequence	other
18774	6 0446826953	F09609	ESTs	7
36722	6,0172343991	AA435512	ESTa	SS.
18062	6.0034342969	AA179845	ESTs Moderately similar to rabionesin-6 (M.musculus)	other

FIGURE 8 (cont.)
13 of 37

22989	5.9992817406	T 16305	EST:	other
41745	5:9905623898	R95895	EST.	7
8787	5.9894877658	AA504307	X-LINKED HELICASE II	other
20550	5.984861795	N55013	EST.	other
26470	5.9417764101	AA262179	ESTs	other
16574	5.9356497569	AA031926	EST	other
693	5,9169537365	030007	Human mRNA for KIAA0165 gene partial cds	other
4093	5.914830973	U25182	Human antioxidant enzyme AOE37-2 mRNA complete	TM
1192	5.9086264407	HG3546-HT3744	EST - HG3546-HT3744	7
22956	5.8954735623	T10248	ESTa	other
36723	5.891608409	AA435524	EST	?
2114	5.6844986595	L40384	E5T - L40384	other
26872	5.858238789	AA291137	ESTs	other
6602	5,8663883018	X98266	EST - X98266_cds2	other
42701	5.8594493433	Z38612	ESTs	other
28573	5.84591116	C21118	ESTs	other
18290	5.6189427595	AA211901	ESTs	other
732	5.8043917941	D83781	Human mRNA for KIAAD197 gene partial cds	other
5330	5.8014145611	U91327	EST - U91327	7
33503	5.7990715189	W88720	EST	7
2553	5.7797505864	M26167	Human ptatelet fector 4 varation 1 (PF4var1) gene	7
34705	5 7658806254	AA286907	complete cds ESTs Weakly similar to pusative p150 [H.sepiens]	other
42665	5.7694091043	W93659	ESTs	other
38180	5.7539310793	AA487495	EST - RC_AA487495	other
4244	5.7476738809	U33286	Human chromosome segregation gene homolog CAS	other
32822	5 7418957453	W16834	mRNA complete cds ESTs	тм
3977	5.7245885557	U18991	Retinal pigment epithelium-specific protein (65kD)	7
24673	5.7202366155	239301	EST	TM
6928	5,7120261128	746629	SRY (sex-determining region Y)-box 9 (campomelic	other
38726	5 7030796758	AA608733	dysplasia autosomel sex-reversal) ESTs	7
39290	5.6892372058	C14573	Human mRNA for KIAA0029 gene partial cds	other
11405	5.6818873796	AA232231	FSTs	other
22538	5.6792008591	R73567	Homo sepiens metrin-L precursor (ADAM12) mRNA	TM
			complete ods	TM
40747	5.6605393208	N56872	Homo sapiens cone 22 mRNA atternative splice variant alpha-1 complets cds	
31596	5.6554024604	N72094	EST:	other
6329	5.6415652518	X82279	EST - X82279	7
31578	5.6273323661	N71361	ESTs ·	other
33207	5 6271818482	W70051	H.sapieris mRNA for M-phase phosphoprotein mpp9	other
2545	5.6105860146	M25753	Cyclin B1	other
22580	5.5988402647	R79156	EST:	other
33592	5.5935314518	W93127	ESTs.	other
28843	5.5734698755	D60252 ·	EST:	plher
6160	5.5689050619	X74794	CDC21 HOMOLOG	other
37967	5.551345667	AA479666	EST4	other
42515	5.5217968611	W72116	Home sepiens clone 23622 mRNA sequence	other
4732	5 5130668527	U58522	Human huntingtin interacting protein (HIPZ) mRNA complete cits	other
3299	5.5099850678	M95623	Hydroxymethy®iane synthase	,
28320	5.473406981	AA599574	EST:	7
746	5,471260899	D84454	Human mRNA for UDP-galactose translocator complete cds	TM
39373	5.4635804954	C21517	EST:	other
3117	5.4396413537	M81182	Peroxisomal membrane protein 1 (70kD Zaflwager syndrome)	other

FIGURE 8 (cont.)
14 of 37

21257	5.4343612441	R0919G	ESTs Moderately similar to M-phase phosphoprotein 11 [H.sapiens]	other
31487	5 4318648859	N69507	ESTs	other
28954	5 4137130511	F03153	ESTs	other
38928	5.389762721	AA609595	ESTs	other
29903	5 3722320622	N23366	EST	7
30925	5 3437432315	N58295	ESTs Weakly similar to LINE-1 REVERSE	,
19091	5 3344615669	H07864	TRANSCRIPTASE HOMOLOG [H.saplens] ESTs	TM
25209	5 3138951918	AA491250	ESTs	other
9470	5.3118897984	H46617	EST - H46617	other
9435	5,3070056656	H30201	EST - H30201	7
28552	5 2954432572	Ç20914	ESTs	other
27411	5.2940164267	AA428137	ESTs	other
30615	5.2924125264	N50556	EST:	cther
26313	5.2657977167	AA599309	ESTs	TM
39321	5.2649035384	C20632	ESTs	7
29934	5.2531047395	N24184	EST.	other
1094	5.2496703122	HG2846-HT2983	EST - HG2646-HT2983	7
39578	5.2461126384	F08925	EST ₆	TM
11232	5 2466798424	AA186804	ESTs Weekly similar to unknown (S.cerevisiae)	other
2466	5.2426349328	M21539	Human small proline nich protein (sprtl) mRNA clone	other
26843	5 2387758661	AA287450	1292 ESTa	7
40331	5,2353385567	H97562	ESTs Weakly smilar to SPERMATID-SPECIFIC	other
6035	5.205798365	AA305116	PROTEIN T2 (Sepia officinalis) EST - AA305116	other
29793	5 1855425722	N20593	ESTs Weakly similar to weak similarity to procollagen	other
20.00			elpha chain 1(V) chain (C.elegens)	••••
34109	5.1481590107	AA210722	EST	7
26408	5.1432577257	AA258177	ESTs Weakly similar to ROSA28AS [M.musculus]	other
19263	5 1427029807	H15054	EST:	ТМ
24596	5.1416089352	238810	E5Ts	other
28589	5.1365059753	C21245 X17098	H.sapiens mRNA for apoptosis specific protein	other
5684	5.1121931412		Pregnancy-specific beta-1 glycoprotein 6	other
30710	5.1079347344	N51761	EST	,
35765	5.0973514948	AA406167 AA256460	EST ST.	7
26360			ESTs .	
2351	5 0849612092	M15796	Proliferating cell nuclear antigen	?
30267	5 0836877534	N35065 T03886	Homo saciens clone 24739 mRNA sequence	other
41792 36710	5.0737512465 5.0703839864	AA434411	ESTs .	,
•			••••	other
39090	5.0546885407	AA620628 779951	EST:	TM
42185 18745	5,0539926381	F09134	ESTs ESTs	? other
35746	5.0396841996	AA406063	ESTs	
35356	5.0354809581	AA399053	EST	other ?
36769	5.0312706878	AA435750	EST	,
36900	5.0312108678	AA436866	H.sapiens mRNA for M-phase phosphoprotein mpp9	other
27595	5 0244757301	AA443326	ESTs	TM
16290	5.0056611904	AA015145	ESTs	7
27117	5 0018148599	AA405098	ESTs Weakly similar to MOESIN/EZRIN/RADIXIN	cther
			HOMOLOG (D.melznogaster)	
4304	4.9951954397	U36764	Eukeryotic translation instation factor 3 (eIF-3) p36 subunit	other
33458	4.9907402071	W86835	Homo sapiens mRNA for KIAA0636 protein complete	other
26693	4.9800090679	AA282120	EST	,
12669	4 9758138651	AA417030	Homo sapiens protein regulating cytokinesis 1 (PRC1)	other

FIGURE 8 (cont.)
15 of 37

	•				
25	701	4 9708526387	H97970	EST	7
20	M80	4.9557253636	N52168	EST:	TM
8	720	4 9439110602	AA481218	EST - AA481218	other
34	1828	4 9431269475	AA292436	Homo sapiens semaphorin F homolog mRNA complete	SS.TM
14	1985	4.941621032	U15128	cds Human beta-12-N-acetylglucosaminyttransferase II (MGAT2) gene complete cds	?
18	3115	4.9377553522	AA004420	EST:	7
42	2506	4.9348587118	W70074	EST	other
34	1761	4 9316837445	AA287633	ESTs :	other
11	870	4 9281056201	AA262587	ESTs .	TM
23	1211	4.9258391854	T40889	ESTs	other
40	611	4.9160502275	N39138	Homo sapiens mRNA for KIAA0584 protein partial cds	other
42	611	4 9128605354	W87006	Homo sepiens putative RNA binding protein KOC (koc) mRNA complete cds	other
39	652	4.9045174605	нозо99	ESTs Companie cos	other
17	581	4,889674751	AA129395	EST	7
37	239	4.8704375389	AA449121	EST.	?
18	712	4.6703618781	F04677	EST.	other
30	1709	4.8611171953	N51752	ESTs Weakly similar to synapse-associated protein sap47-1 [D.metenogaster]	other
34	179	4.8503613948	AA227903	ESTs Highly similar to GTP-BINDING PROTEIN LEPA {Pseudomonas fluorescens}	other
21	433	4.625670988	R22183 .	EST	?
39	731	4.6186142741	H11760	EST6	other
31	295	4.8116614607	N66653	EST.	other
24	647	4.804163055	239106	EST	?
31	292	4:6006871817	N66515	EST\$	other
12	265	4.7997542393	HG4157-HT4427	EST - HG4157-HT4427	7
1	106	4 7932425858	HG2981-HT3127	EST - HG2981-HT3127	7
18	212	4.7912262565	AA196506	ESTs	other
34	367	4.782207045	AA251758	Homo sapiene spleen mitolic checkpoint BUB3 (BUB3) mRNA complete cds	other
34	802	4.7797760205	AA291468	ESTs	TM
34	762	4.7775301546	AA287834	EST:	other
11	595	4.7696612848	AA242819	EST#	other
83	295	4,7639839111	AA405082	ESTs	7
17	622	4.758635576	AA131584	ESTs Weakly similar to SOF1 PROTEIN	other
35	781	4.7572463523	AA406335	ESTs	other
34	754	4.7483874972	AA287642	Human mRNA for KIAA0078 gans complete cds	other
23	237	4.7444854356	T47291	EST	7
37	667	4.7280445357	AA460318	ESTS Highly similar to 60S RIBOSOMAL PROTEIN L26 [Rattus norvegicus]	other
	568	4.7257189975	AA236786	EST:	other
38	622	4.7190695733	AA598967	EST:	?
	137	4.7057359474	U79296	Dihydrolipoamide S-acetyltransferase (E2 component of pyruvale dehydrogenase complex)	other
	038	4.7002244728	AAD10065	CDC26 protein kinase 2	other
	288	4.7000147312	H16567	ESTs	other
	503	4.6979488292	T17045	Collagen type I stpha-2	other
-	278 396	4.6953739298	M94055 £38961	SODRUM CHANNEL PROTEIN BRAIN II ALPHA SUBUNIT Integral transmembrane protein 5	TM TM
-	400	4,6901390898	AA399591	Homo espiens putative DNA methyltransferase	other
-	246	4.6662691303	AA396367	(DNMT2) mRNA complete cds EST Weakly similar to HSP60 protein [M.musculus]	7
36	387	4.6822499271	AA426270	ESTs	other
	509	4.6730072542	R27314	ESTs	olher
	381	4.6729672124	N67889	ESTs	other
26	723	4 6727894925	AA282781	ESTs Highly similar to PRE-MRNA SPLICING FACTOR	othe/

FIGURE 8 (cont.)
16 of 37

	4 6703621086	AA425151	Human GAP SH3 binding protein mRNA complete cds	other
17409	4 6688418667	AA113136	EST - RC_AA113136	other
4908	4.6552339935	U67156	Human mitogen-activeted kinase kinase kinase 5 (MAPKKKS) mRNA complete cds	other
30594	4.6496238328	N49967	ESTs	other
38286	4.64639735	AA489847	ESTs Weakly similar to LINE-1 REVERSE TRANSCRIPTASE HOMOLOG (Nyclosbus coucang)	η.
13073	4 6426509459	AA433950	ESTs	other
40435	4.6240161066	N21614	Homo sapiens basic-leucine zipper transcription factor MatG (MAFG) mRNA complete cds	other
14474	4.6228694379	AA609427	ESTs Moderately similar to IIII ALU SUBFAMILY SC WARNING ENTRY IIII (H. sapions)	other
38213	4.815309907	AA488847	ESTs Weakly similar to putative p150 [H.sapiens]	7
5312	4.606644198	U90716	Human cell surface protein HCAR mRNA complete cds	55.TM
24225	4,6041550359	W70326	ESTs	7
35588	4.5868982366	AA401750	EST	7
29739	4,5863199051	H99626	EST .	7
7203	4,5792992577	AA053096	EST - AA053096	other
2157	4.5772055869	L41939	Homo sepiens protein-tyrosine kinese EPHB2v (EPHB2) mRNA complete ods	
32086	4.5661024279	R11510	EST:	7
8085	4.5648114738	AA314779 D13633	ESTs Weakly simitar to LITHOSTATHINE 1 BETA PRECURSOR (H. sapiens) Human mRNA for KIAA0008 gene complete cds	5S. other
224				other
34006	4.5609980241	AA188761	DNA polymerese gamma	
33656	4.5557384389	W95477	ESTs	other
34065	4.5537335124	AA195517	ESTs Weakly similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII [H.sapiens]	TM
6026	4.5357922097	X66503	Adenylosuccinale synthase	other
4165	4.5032930671	U29463	Cytochrome 8561	7
40262	4.5024727522	H93562	ESTs	TM
22687	4.5018672549	R88209	ESTs	TM
41069	4.4977510482	N93969	H.sapiers mRNA for hFat protein	S\$.
8264	4.4793100575	AA401334	ESTS	other
27588				
2/300	4.472017297	AA443187	ESTs	other
35882	4.472017297 4.4717597552	AA443187 AA412047	ESTs ESTs	other 7
35882 34479	4.4717597552 4.465519191	AA412047 AA262080	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCC1) mRNA complete cds	7 TM
35882 34479 15921	4.4717597552 4.485519191 4.4548516436	AA412047 AA262080 Y12065	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCC1) mRNA complete cds Homo sepiens mRNA for nucleolar protein hNop56	7 TM ?
35882 34479 15921 11279	4.4717597552 4.485519191 4.4548516436 4.4380038671	AA412047 AA262080 Y12065 AA195399	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRNA complete cds Homo septems mRNA for nucleolar protein hNop56 ESTs	7 TM ? other
35882 34479 15921	4.4717597552 4.485519191 4.4548516436	AA412047 AA262080 Y12065	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCC1) mRNA complete cds Homo septens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Saccharomyoas	7 TM ?
35882 34479 15921 11279	4.4717597552 4.485519191 4.4548516436 4.4380038671	AA412047 AA262080 Y12065 AA195399	ESTs Human burnetanide-sensitive Na-K-CI colransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA-	7 TM ? other
35882 34479 15921 11279 39222	4.4717597552 4.465519191 4.4548516436 4.4380038671 4.4367650786	AA412047 AA262080 Y12065 AA195399 AA621348	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- CLUCOSYLTRANSFERASE [Saccharomyoes cerevisiae]	7 TM ? other other
35882 34479 15921 11279 39222 34428	4.4717597552 4.465519191 4.4548516436 4.4380038671 4.4367650766 4.4364736766	AAA12047 AA262080 Y12065 AA195399 AA621348	ESTs Human burnetanide-sensitive Na-K-Cl cotransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Saccharomyces corviviate] ESTs Homo sepiens mRNA from chromosome 5q21-22	7 TM ? other other
35882 34479 15921 11279 39222 34426 6771 22193 7698	4.4717597552 4.465519191 4.4548516436 4.430038671 4.4367650786 4.4364736766 4.43267373 4.4189610024 4.4066170674	AA412047 AA262080 Y12065 AA495399 AA621348 AA256526 AA491188 R53891 AA263032	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleoler protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Seccharomyoes cerevisiae] ESTs Homo sepiens mRNA from chromosome 5q21-22 clone.A3-A ESTs	7 TM 7 other other other other
35882 34479 15921 11279 39222 34428 8771 22193	4.4717597552 4.465519191 4.4548516436 4.4380038671 4.4367650786 4.4364736766 4.43267373 4.4189610024	AA412047 AA252080 Y12065 AA195399 AA621348 AA258526 AA491188 R53891	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRNA complete ds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Saccharomyoes cerevisiae] ESTs ESTs ESTs Lono-SAS-A Homo sapiens mRNA from chromosome 5q21-22 chone-AS-A	7 TM ? other other other other
35882 34479 15921 11279 39222 34426 6771 22193 7698	4.4717597552 4.465519191 4.4548516436 4.430038671 4.4367650786 4.4364736766 4.43267373 4.4189610024 4.4066170674	AA412047 AA262080 Y12065 AA495399 AA621348 AA256526 AA491188 R53891 AA263032	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleoler protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Seccharomyoes cerevisiae] ESTs Homo sepiens mRNA from chromosome 5q21-22 clone.A3-A ESTs	7 TM 7 other other other other
35882 34479 15921 11279 39222 34428 8771 22193 7698 19902	4.4717597552 4.465519191 4.4548516436 4.430033671 4.4367650786 4.4364738768 4.432067373 4.4189610024 4.4066170674 4.3866145805	AA412047 AA262080 Y12065 AA495399 AA621348 AA256526 AA491188 R53891 AA263032 H66736	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Saccharomyoes cerevisiae] ESTs Homo sepiens mRNA from chromosome 5q21-22 clone-A3-A ESTs ESTs ESTs ESTs	7 TM 7 other other other other other other other
35882 34479 15921 11279 39222 34428 6771 22193 7698 19902 9276	4.4717597552 4.465519191 4.4548516436 4.4380038671 4.4367650786 4.4364736769 4.43267373 4.4189610024 4.4066170674 4.3866145805 4.3866095209	AA412047 AA262080 Y12065 AA195399 AA621348 AA256526 AA491188 R53891 AA263032 H66736 D82374	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Saccharomyoes cerevisae] ESTs	7 TM ? other other other other other other other
35882 34479 15921 11279 39222 34428 8771 22193 7698 19902 9275 10716	4.4717597552 4.465519191 4.4548516436 4.4380038671 4.4367650786 4.4364738768 4.432067373 4.4189610024 4.4066170674 4.3866145805 4.386605209 4.3784529068	AA412047 AA262080 Y12065 AA195339 AA621348 AA256526 AA491188 R53891 AA263032 H66736 De2374 AA053319	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Saccharomyoes cerevisiae] ESTs Homo sepiens mRNA from chromosome 5q21-22 clone-A3-A ESTs ESTs ESTs ESTs	7 TM 7 other other other other other other other
35882 34479 15921 11279 39222 34428 8771 22193 7698 19902 9276 10716 13193	4.4717597552 4.465519191 4.4548516436 4.4380038671 4.4367650786 4.43267373 4.4189610024 4.4066170674 4.3866145805 4.3868095209 4.3751913512	AA412047 AA262080 Y12065 AA195339 AA621348 AA256526 AA491188 R53891 AA263032 H66736 De2374 AA053319 AA412763	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Saccharomyoes cerevisiae] ESTs Homo sepiens mRNA from chromosome 5q21-22 clone-A-A- ESTs	7 TM 7 other other other other other other other other
35982 34479 15921 11279 39222 34426 8771 22193 7698 19902 9276 10716 13193 5690	4.4717597552 4.485519191 4.4548516436 4.438038671 4.4367650786 4.4364736768 4.43267373 4.4195610024 4.4066170674 4.3866145805 4.3866145805 4.3761913512 4.3751913512	AA412047 AA262080 Y12065 AA195399 AA621348 AA256526 AA491188 R53891 AA283032 H66736 D82374 AA053319 AA442763 X17620	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Saccharomyoes cerevisiae] ESTs Homo sepiens mRNA from chromosome 5q21-22 clone-A3-A ESTs ESTS	7 TM 7 other
35982 34479 15921 11279 39222 34428 8771 22193 7698 19902 9276 10716 13193 5690 35102	4.4717597552 4.465519191 4.4548516436 4.438038671 4.4367650786 4.432067373 4.4189610024 4.4066170674 4.3866145805 4.3866145805 4.3761913512 4.3751913512 4.3723059417 4.37147138	AM12047 AA262080 Y12065 AA195399 AA621348 AA258526 AA491188 R53891 AA683032 H66736 De2374 AA053319 AA42763 X17620 AA371509 AA169228 AFEX.	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Saccharomyces orrorisiae] ESTs Homo sepiens mRNA from chromosome 5q21-22 clone-AS-A ESTs ESTS	7 TM ? other other other other other other other other other TM other
35882 34479 15921 11279 39222 34426 6771 22193 7698 19902 9276 10716 13193 5690 35102 17983	4.4717597552 4.465519191 4.4548516436 4.436038671 4.4367650786 4.432067373 4.4189610024 4.4066170674 4.3866145805 4.3764529068 4.3761913612 4.3723059417 4.37147138 4.3612985467	AM12047 AA262080 Y12065 AA195399 AA621348 AA256526 AA491188 R53891 AA263032 H66736 D82374 AA653319 AA42763 X17620 AA371509 AA169228 AFEX.	ESTs Human burnetanide-sensitive Na-K-Cl cotransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Saccharomyoes corevisase] ESTs Homo sepiens mRNA from chromosome 5q21-22 clone-A3-A ESTs	7 TM ? other other other other other other other the other TM other
35882 34479 15921 11279 39222 34428 8771 22193 7698 19902 9276 10716 13193 5690 35102 17983 24962	4.4717597552 4.465519191 4.4548516436 4.436038671 4.4367650786 4.43267373 4.4189610024 4.4066170674 4.3866145805 4.37647352068 4.3751913612 4.3723059417 4.37147138 4.3612985467 4.3497206925	AA412047 AA262080 Y12065 AA1953399 AA621348 AA256526 AAA91188 R53891 AA263032 H66736 D82374 AA053319 AA442763 X17620 AA371509 AA169228 AFFX. HUMTFRR/M115 07	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRIAA complete cds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE (Seccharomyoes cerevisiae) ESTs ESTs Homo sepiens mRNA from chromosome 5q21-22 clone-JA-A ESTs EST- RC_AAJ71509 ESTs AFFX-HUMTERRAM11507_5	7 TM 2 other other other other other other other other the other TM other other TM other 2
35882 34479 15921 11279 39222 34428 8771 22193 7698 19902 9276 10716 13193 5690 35102 17983 24962	4.4717597552 4.465519191 4.4546516436 4.436038671 4.4367650786 4.43267373 4.4189610024 4.4066170674 4.3866145805 4.3868095209 4.37541913512 4.3723059417 4.37147138 4.3612985467 4.3497205925 4.3418539669	AA412047 AA262080 Y12065 AA195399 AA621348 AA256526 AA491188 R53891 AA263032 H66736 D62374 AA053319 AA412763 X17620 AA371509 AA169226 AFEX. HUMTERPARTIS TO TO T	ESTs Human burnetanide-sensitive Na-K-Cl colransporter (NKCCI) mRNA complete cds Homo sepiens mRNA for nucleolar protein hNop56 ESTs ESTs Highly similar to DOLICHYL-PHOSPHATE BETA- GLUCOSYLTRANSFERASE [Seccharomyoes cerevisiae] ESTs Homo sepiens mRNA from chromosome 5q21-22 clone-JA-A ESTs	? TM ? other other other other other other other other other TM other other TM other other ?

FIGURE 8 (cont.)
17 of 37

16335	4 3019961487	AA018587	ESTs Weakly similar to !!!! ALU SUBFAMILY SP WARNING ENTRY !!! (H.sapiens)	7
33036	4.2915644973	W48560	ESTs Weakly similar to transposon LRE2 reverse transcriptase homolog [H.sapiens]	other
30180	4.2697721925	N33144	ESTs	other
35591	4.2895541242	AA401758	ESTs Wealdy similar to IIII ALU SUBFAMILY SQ WARNING ENTRY IIII (H.sapiens)	SS.
25340	4.2721717135	AA054554	EST .	?
28106	4.2659103748	AA485084	ESTs	other
38690	4.2649184307	AA600121	ESTs	other
20203	4.2626499431	N26855	ESTs Moderately similar to IIII ALU SUBFAMILY SQ WARNING ENTRY IIII [H.sapiens]	other
10251	4.2608760694	R76185	ESTs Weakly similar to C01H6.7 [C.elegans]	SS.
12684	4.2604192389	AA417558	EST&	5 \$.
31636	4.2509469427	N73680	Natural resistance-essociated macrophage protein 2	TM
20769	4.2479765348	N67277	ESTs .	19/20
1572	4.2353281083	K01884	EST - K01894	7
10923	4.2292322072	AA116036	EST•	other
34380	4.2283792392	AA252414	EST6	other
10132	4.2222816115	R35733	EST - R35733	other
16629	4.2161752119	AA036811	ESTs	other
25146	4.1969683794	AA026356	ESY:	7
28730	4.1965943098	020959	ESTs Moderately similar to IIII ALU SUBFAMILY SQ	other
10200	4.1874912391	R64521	WARNING ENTRY IIII [H.sapiens] ESTs	other
38695	4.1545794663	AA600176	ESTs	other
31365	4.150549979	N67550	ESTs	other
42379	4.1496120668	W37999	EST#	other
28050	4.1428703354	AA479139	Acid phosphatase 1 soluble	ather
2620	4.1386565707	M29474	Human recombination activating protein (RAG-1) gene	,
8927	4.1340593744	AF008442	complete cds Homo sapiens RNA polymerase i subunit hRPA39 mRNA complete cds	other
13379	4.1269549188	AA449741	ESTs Weakly similar to AF-9 PROTEIN [H.sapiens]	other
5134	4.1218251608	U79293	Human clone 23948 mRNA sequence	other
2626	4.1213948	M29581	Zinc finger protein 6 (clone HF.16)	other
38005	4.1160483668	AA479969	ESTS	other
36575	4.1127196584	AA431085	EST	,
18296	4.1121837207	AA213620	ESTe Weakly similar to putative p150 (H.sapiens)	,
29531	4.1111459313	H88953	EST - RC_H88953	TM
143	4.1095880506	AFFX- HUMTFRRM115	AFFX-HUMTERRAM11507_5	7
10970	4.0967613396	07 AA129390	ESTe	other
25836	4.0952825397	- AA152305	Interferon (gamma)-induced cell line protein 10 from	\$8.
19735	4.0937927853	H53038	EST	,
40711	4.0909709431	N53564	ESTE	other
4149	4.0901471427	U28386	RAG (recombination activating gene) cohort 1	TM
5767	4.0862784557	X53793	MULTIFUNCTIONAL PROTEIN ADE2	other
5503	4.0861035825	X05232	Stromelysin	SS.
20310	4.0641711656	N34893	ESTs Highly similar to HYPOTHETICAL 47.8 KD PROTEIN 80280.9 IN CHROMOSOME III [Caenorhabditis electors]	other
458	4.0599824566	D38145	Prostaglandin I2 (prostacyclin) synthase	SS,
7614	4.0\$59685576	AA248406	ESTs	other
40230	4.0447282719	H90181	EST»	88,
33651	4.039204804	W95409	ESTs	other
16777	4.0231657929	AA046968	EST .	7
19110	4.0094905222	H06778	ESTs	other
34442	4.0077010365	AA258093	HKR-T1	other
5099	4 004992433	U79247	Human clone 23599 mRNA sequence	TM

FIGURE 8 (cont.)
18 of 37

6209 .	3.9990473163	AA384220	ESTs ·	other
24408	3.9976586074	W90146	£5Ts	other
26596	3.9974919787	AA279943	EST»	other
16485	3.9811264008	AA026269	Spleen facus forming virus (SFFV) provins integration oncogene spi1	other
32969	3.9804901745	W42451	ESTs	TM
27006	3.9799768093	AA398695	ESTs Wealthy similar to E04F6.2 game product	other
29809	3.9526765967	N21043	(C.elegans) EST .	7
9596	3.9440163451	H91564	ESTs .	TM
29024	3.9377933938	F09315	Homo expisins mRNA for KIAA0583 protein partial ods	ather
21694	3.9356365584	R39317	Homo sapiens protein-tyrosine kinase EPH82v (EPH82)	other
13207	3,929998104	AA443321	mRNA complete cds ESTs	other
37865	3 9143752629	AA476623		pther
3/003	3 01437 02525	701.70020	ESTs Highly similar to PUTATIVE UBIQUITIN CARBOXYL-TERMINAL HYDROLASE C13A11.04C [Schizosaccharomycas pombe]	
36201	3.8129828172	AA421164	EST	7
6961	3 898 1 160269	AFFX- HUMTFRRAM115	AFFX-HUMTFRRM11507_3	7
17444	3.8927133917	07 AA115933	EST.	other
25869	3.8919834527	AA157267	ESTS Highly similar to HYPOTHETICAL 37.7 KD PROTEIN ZKESS.3 IN CHROMOSOME III	TM
24852	3.89042252	Z41415	[Ceenorhabditis elegans] ESTs Highly similar to BONE MORPHOGENETIC	other
26685	3.689363206	AA281950	PROTEIN 1 PRECURSOR [Mus musculus] ESTs	7
42300	3.6850230366	T95850	ESTs	7
6495	3.8830844863	X92715	Zinc finger protein 74 (Cos52)	other
38604	3.8828045942	AA598803	ESTs	TM
36358	3.8826713718	AA425756	ESTs	other
30560	3 873276445	N49264	MYB PROTO-ONCOGENE PROTEIN .	other
14413	3.8724466158	AA600150	ESTs	other
23823	3.8574824967	T91805	Homo sepiens mRNA for SY1C2 complete cds	other
38158	3.853096838	AA487021	EST	?
2572	3 8519747554	M27281	Vascular endothelial growth factor	other
40100	3.8484168967	H75933	Laminin receptor (2H5 epitope)	other
40258	3.8462992993	H93340	EST:	TM
20944	3.8461621525	N74443	ESTs	other
20411	3.8459400966	N48953	Homo sapiens mRNA for KIAAC689 protein portial cds	other
10345	3.8457714481	AA001663	EST:	other
31261	3.8451974374	N66248	EST	other
8513	3.8376410994	AA446990	EST:	other
13877	3.8363409835	AA476604	ESTs .	other
40748	3.8253562321	N56879	EST	7
14509	3.8152852193	AA609943	EST:	other
10281	3.8065567331	R80333	EST	other
25284	3.8044158642	AA045074	ESTs Weekly similar to 52-kD SS-A/Ro autoantigen {H.saptens}	other
6730	3.7900025129	Y09305 .	H.sapiens mRNA for protein kinase Dyrk4 partial	other
16033	3.7684592402	AFFX- HUMISGF3AM9 7935	AFFX-HUMISGF3A/M97935_MB	,
39242	3.7627164808	AA621523	EST ₃	other
27354	3.7794760435	AA425221	EST ₈	7
4552	3,777263605	U49168	Human placente (Diff33) mRNA complete cds	\$8.TM
18365	3.7756199108	AA227219	Homo sapiens CAGF9 mRNA partial cds	other
16754	3.7677416053	AAD460G7	EST - RC_AA046067	other
12752	3.7671137403	AA421250	ESTs	other
42463	3.7601033108	W50180	ESTs	other

FIGURE 8 (cont.)
19 of 37

10614	3 7581669016	AA037357	ESTs	7
867	3.7459337969	D87716	Human mRNA for KIAA0007 gene partial cds	other
7608	3 7236047135	AA180967	ESTs	other
31785	3.732736742	N80703	EST:	other
35377	3 7273784503	AA399453	EST - RC_AA399453	7
22828	3 7243928524	R98192	ESTs	other
25240	3.7243198336	AA039713	EST:	other
11008	3.7197361366	AA134289	ESTs Weakly similar to ASH1 [D.melanogaster]	7
4341	3.7162349944	U38545	Human ARF-activated phosphatidylcholine-specific phospholipase D1a (hPLD1) mRNA complete cds	ather
28833	3.7147818393	D59787	EST - RC_D59787_1	7
3750	3.7121007154	U09279	Cottagen type XIX alpha 1	85.
17483	3 6943413512	AA122147	EST•	TM
16854	3 6915208471	AAD55552	ESTs Weakly similar to KIAA0319 [H.sapiens]	TM
3709	3.6891656771	U07550	Heat shock 10 kD protein 1 (chaperonin 10)	other
1608	3.6652978422	L00205 '	KERATIN TYPE II CYTOSKELETAL 6D	, ,
24577	3.6617721053	Z38727	Homo sepiens mRNA for KIAA0555 protein complete ods	TM
31032	3.6570916388	N62508	EST:	other
4951	3.6538195433	U69546	Human RNA binding protein Etr-3 mRNA complete cds	nertta
37660	3,6523275307	AA460225	EST4	other
20418	3.6495357091	N49209	EST:	other
27995	3 6485167436	AA470155	Homo sapiens coatomer protein (COPA) mRNA complete ods	7
7971	3.6434397105	AA287423	EST#	other
27606	3.84303453	AA443793	EST:	other
24677	3,6427250633	239338 AA148521	ESTs Highly similar to POLIOVIRUS RECEPTOR HOMOLOG PRECURSOR (Mus musculus) ESTs Weakly similar to putative p150 [H.sapiens]	other TM
11070	3.6356048599	D89618		
9328 36826	3.6356048599	AA435998	Home sapiens importin-alpha hometog (SRP1gamma) mRNA complete ods ESTs	other
17678	3.6300045795	AA134275	Human HIV1 tata element modulatory factor mRNA	other
36209	3 6274694477	AA421266	sequence from chromosome 3 ESTs Wealdy similar to LIS-1 protein [H.sapiens]	olner
34120	3.6258090412	AA211615	EST	7
38152	3.6248442011	AA486737	H.sapiens mRNA for Sm protein F	TM
38463	3.6164693268	AAS04491	ESTs Weakly similar to contains similarity to C3HC4-	TM
		•	class zinc finger (C.alegans)	
20064	3.6183699978	H98653 N66152	ESTs	тм
31256	3.5992620732		EST	7
9713	3.5985226843	L44338	Homo sapiens mRNA for KIAA0525 protein partial cds	other
28622	3.5768056147	D11837	ESTs	7
38057	3.5738105703	AA481549	EST - RC_AA481549	other
26763	3.5688723791	D45568	EST	7
16996	3.5680705709	AA069038	EST - RC_AA069038	TM
28628	3.5604144617	D11888	ESTs Moderately similar to PROHIBITIN (H.sapiens)	7
25804	3.5442954572	AA148885	ESTs	?
2492	3,5423964239	M22898	Tumor protein p53 (Li-Fraumani syndrome)	7
14904	3.5411970737	TB3389	ESTs Highly similar to GEPHYRIN (Rattus norvegicus)	other
25265	3.5347588502	AA043765	H.sapiens RY-1 mRNA for putative nucleic ecid binding protein	ather
13606	3.5327912417	AA456437	ESTs Weekly similar to CLEAVAGE STIMULATION FACTOR 64 KD SUBUNIT [H.sapiens]	other
42307	3.5318436465	T96595	EST - RC_T96595	TM
1544	3.526202414	J05068	TRANSCOBALAMIN I PRECURSOR	88,
42339 42311	3.5195061035 3.5183719631	W02072 T97257	ESTs Weakly similar to No definition line found [C.alegans] ESTs	other
	3 5183/19631	197257		
2023	3.5040279423	1.34600	INITIATION FACTOR IF-2 MITOCHONDRIAL PRECURSOR	other

FIGURE 8 (cont.) 20 of 37

4540	3.4955308569	U48807	Human MAP kinase phosphatase (MKP-2) mRNA complete cds	other
33707	3.4888534277	Z3929 7	Neuronal pentration II	other
17220	3 4755763461	AA083070	EST - RC_AA083070_s	SS.
24332	3.4725273806	W85782	ESTs	otner
35887	3 4668063718	AA412067	ESTs .	other
20158	3.4538150055	N23638	ESTs Weakly similar to coded for by C, elogans cDNA yk52e10.5 [C.elegans]	other
8338	3 4465832071	AA417152	Home samens protein regulating cytokinesis 1 (PRC1)	other
387	3.4421827234	D28589	mRNA complete cds EST - D28589	other
12319	3.4356269717	AA398109	ESTs .	6S,TM
38276	3.4313139432	AA489711	EST:	TM
15643	3,4312194246	W58247	ESTs Highly similar to KINESIN-LIKE PROTEIN KIF4	ciher
11218	3.4232932843	AA180488	[Mus musculus] ESTs	TM
16539	3.417686379	AA029328	Human mRNA for KIAA0073 gene partial ods	,
29703	3.4162847487	H28581	EST:	other
13838	3.4162403464	AA465342	EST:	other
25585	3.4160353003	AA112389	H4(D10S170)	S S,
34018	3 4145338583	AA191488	Human high-effinity copper uptake protein (hCTR1) mRNA complete cds	TM
251	3.4006042651	D14520	Basic transcription element binding protein 2	Other
3778	3.4004516201	U09849	Zinc tinger protein 139 (clone pHZ-37)	other
24535	3.3984397637	238409	ESTs	other
16858	3.3925194041	AA055759	Human mRNA for KIAA0128 gene partial cds	тм
16127	3.3921645927	AA004569	ESTs	other
36683	3.3841316491	AA432268	EST.	other
26149	3.3809497785	AA250624	ESTs Weakly similar to ZINC FINGER PROTEIN 91 (H.sapiens)	other
4011	3.3798093471	U20536	Human cysteine protease Mich2 Isoform alpha (Mich2) mRNA complete cds	other -
41001	3.3794250205	N78844	EST&	other
5660	3.3789336731	X16396	NAD-DEPENDENT METHYLENETETRAHYDROFOLATE DEHYDROGENASE	S5.
19204	3.3778332343	H11629	EST:	other
42323	3.3768515979	T98152	Fibrillan 2	SS .
26928	3.3725378868	AA342580	ESTs	58.
20497	3,369285912	N52565	E\$Ts	other
19726	3.36674249	H12455	EST:	other
36267	3.3606641838	AA424046	ESTs	other
32257	3.3559796018	R54726	DNA-REPAIR PROTEIN XRCC1	other
17365	3.3522214732	AA101551	EST ₈	other
15296	3.3491193196	W16684	ESTs Moderately similar to Similar to Sicerevisies hypothetical protein L3111 (H.sapiens)	other
17675	3.3485870272	AA134064	EST:	TM
40332	3.3456469589	H97565	Home sapiens mRNA from chromosome 5q21-22 clone:A3-A	Other
7219	3.3385684843	AA056319	Homo sapiens protein phosphatase 2A 856-epsilon (PP2A) mRNA complete cos	other
10006	3.3327827922	N81193	Homo sepiens mRNA for KIAAD628 protein complete cds	7
33985	3.3276877441	AA181580	Homo sapiens importin beta subunit mRNA complete	other
9570	3.3263855302	H85169	Home sapiens ecdium/myo-inositol cotransporter (SLCSA3) gene complete cds	other
37551	3.3155406577	AA456679	ESTs	other
886	3.3111782759	D88613	Human mRNA for hGCMa complete cds	other
23650	3.3069426629	T86293	EST&	other
18367	3.3007433533	AA224180	ESTs Moderately similar to overian-specific protein [R.norvegicus]	,
42494	3.2908070546	W59385	H.sapiens NuMA gene (Clone T33)	other
14310	3.2753564661	AA598412	ESTs Highly similar to HYPOTHETICAL TRP-ASP REPEATS CONTAINING PROTEIN IN PMTS-PCT1 INTERGENIC REGION [Seccharomyces corevisies]	SS.TM

REPEATS CONTAINING PROTEIN IN PARTS-PCT INTERGENIC REGION [Succharomycas corevisios]

FIGURE 8 (cont.)

21 of 37

19233	3 274418299	H12634	ESTs	gther
42283	3.2731086284	194343	Home sapens M962 protein spliced isoform 2 mRNA	other
12809	3.271352097	AA42440G	complete cds ESTs	olner
36265	3.2696023617	AA424489	ESTs	other
21555	3.2666296446	R33073	EST	7
13767	3.266\$695616	AA463234	EST:	TM
4739	3.2661591937	U58766	Human FX protein mRNA complete cds	other
7258	3.263106866	AA075427	EST:	other
17041	3.2629042076	AA070384	EST - RC_AA070354	7
15504	3.2616745245	W28362	EST:	other
23793	3.2811829896	T90971	EST - RC_T90971	other
18214	3.2572346955	AA196635	ESTs	TM
7401	3.257164123	AA094800	Human translation initiation factor eIF3 p66 subunit mRNA complete cds	other
18912	3.2553600001	F10913	Homo sapiena clone 23617 unknown mRNA partial cds	other
36317	3.2509495347	AA425069	Human mRNA for KIAA0334 gene complete cds	7
9410	3.2507279851	H20443	H.sapiens mRNA for TRE5	other
2146	3.2464307696	L41390	EST - L41390	7
18683	3.240814336	F04258	ESTs Highly similar to INORGANIC	7
33891	3.2392191408	AFFX- HUMTFRRM115 07	PYROPHOSPHATASE [Bos laurus] AFFX-HUMTFRRM11507_M	?
14435	3.2372161315	AA608730	ESTs Weakly similar to ELONGATION FACTOR 1-	other
9584	3,2363829855	H88126	ALPHA (Giardia intestinate) ESTs Highly similar to GTP-BINDING PROTEIN LEPA	other
22061	3.2340098572	R49216	(Pseudomonas fluorescens) ESTs	TM
35796	3.233287605	AA410223	EST - RC_AA410223	, ·
37403	3,2261652043	AA453613	ESTs	other
15796	3.2260359988	X16889	ALPHA-GALACTOSIDASE A PRECURSOR	SS.
15840	3,2257932439	X70944	PTB-ASSOCIATED SPLICING FACTOR	other
7518	3.2252170427	AA147144 .	EST - AA147144	other
32335	3.2228388982	R78248	EST:	other
3256	3.2180538038	M92439	130 KD LEUCINE-RICH PROTEIN	other
4400	3.2173896081	U41387	Human Gu protein mRNA partial ods	other
7681	3.2074414299	AA206983	Homo sapiens mRNA for DRIM protein	other
15676	3.2041299443	W68849	ESTs ·	TM
39590	3.2038953621	F09281	EST	other
26883	3.1980022253	AA291921	ESTs Weakly similar to putative p150 [H.sepieris]	7
9808	3,1920380384	M80627	Transcription factor 12 (HTF4 helix-loop-helix transcription factors 4)	other
27755	3.1900699454	AA453444	ESTs	other
29953	3.1682280623	N26011 ·	ESTs	7
21350	3.1876957756	R15846	EST:	other
11981	3.1870525747	AA280928	ESTs .	other
23930	3.1817500097	T96690	ESTs Weakly similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII [H.sapiens]	other
30399	3.1792054412	N45226	EST (M. Espens)	7
22286	3.1781990049	R59312	EST:	other
13494	3.1673900969	AA453431	ESTs	TM
12908	3.1530533441	AA427579	ESTs	other
22319	3.1469419301	R60567	ESTs .	TM
31309	3.1466750623	N66818	ESTs.	TM
31192	3.1458779823	N64406	ESTs ·	other
11288	3.144653134	AA196517	ESTs .	TM
170	3.1430726349	D00596	Thymidylate synthase	,
5307	3.1347905628	U90549	Human non-histone chromosomat protein (NHC) mRNA complete cos	other

FIGURE 8 (cont.) 22 of 37

26105	3 1311103325	AA243133	Homo sapiens serine/threonine kinasa (STAK) mRNA	other
		AA251909	complete cds Horno saprens MAD3-like protein kinese mRNA	other
11659	3 1281786108		complete ads	
19177	3 124408565	H10984	ESTs .	TM
8369	3 1241545824	AA425230	ESTs	TM
34087	3 1216555797	AA205125	Protein serine/threonine kinase stk2	other
25001	3.1209327466	AA004718	ESTs Weakly similar to BAP31 protein (H.sapiens)	other
14149	3.1198500308	AA489665	EST.	other
10167	3,1191986923	R55076	ESTs	other
17380	3,1071055868	AA102566	ESTs	other
42397	3 1044680628	W42928	ESTe	other
14935	3 1042015743	T94828	ESTs Weakly similar to GA BINDING PROTEIN BETA-2 CHAIN [H.sapiens]	other
41673	3.1030349819	R78618	ESTs Weakly similar to GTP-binding protein rab10 [R norvegicus]	ciner
2750	3.1026223619	M35999	Integral beta 3 (platelet glycoprotein (ills antigen CD61)	7
3190	3.1026223619	M95808	PYRUVATE DEMYDROGENASE E1 COMPONENT ALPHA SUBUNIT TESTIS-SPECIFIC FORM PRECURSOR	7
17406	3 0999394188	AA112979	Homo sepiens mRNA for VRK1 complete cds	other
598	3.0912414004	D59253	Homo sepiens mRNA for low molecular mass	other
29348	3,0802365759	H69021	ubiquinone-binding protein complete cds ESTs	other
14130	3 0744457534	AA489041	ESTs	other
14134	3 069660341	AA489080	ESTs Highly similar to phosphorylation regulatory	other
42421	3.0684159011	W45491	protein HP-10 (H.expiens) ESTs Weekly similar to T23G11.7 [C.elegans]	other
15723	3.0660746209	W79060	ESTs Highly similar to ribosome-binding protein p34	other
	3.0650815198	AA158132	(R norvegicus) ESTs Highly similar to YSA1 PROTEIN	other
11140	• • • • • • • • • • • • • • • • • • • •		[Saccharomyces cerevistee]	
28531	3.0649767987	C20679	ESTA	other
2021	3.0628707497	L34409	Homo Szpiens (clone 8383E13) chromosome 4p16.3 DNA fregment	,
14522	3.058260163	AA610108	ESTs Highly similar to PROBABLE PEPTIDYL- PROLYL CIS-TRANS ISOMERASE C21E11.05C [Schizosaccharomyces pombe]	S\$.
29853	3.0545821815	N22162	EST:	other
15962	3.0521475703	Z21420	EST:	other
6541	3.0509806038	X95632	Human Abl interactor 2 (Abi-2) mRNA complete cds	other
13229	3 0485366337	AA443811	EST:	other
27315	3,046622812	AA424038	EST:	other
13621	3.0302305369	AA456821	EST& Highly similar to BONE MORPHOGENETIC PROTEIN 1 PRECURSOR (Mus musculus)	other
35929	3.0269182409	AA412429	EST:	other
17925	3.0253428426	AA164209	Homo sapiens RRM RNA binding protein Gry-rbp (GRY- R8P) mRNA complete cds	other
5053	3 0249536782	U76992	Human Tat-SF1 mRNA complete cds	other
15060	3.0213293848	U54999	Human LGN protein mRNA complete cds	other
17757	3 0205801351	AA147224	EST	?
19050	3.0192379314	H05509	ESTs .	other
26530	3.0176823278	AA278650	ESTs	other
16806	3.0158779932	AA053258	Homo sepiens mRNA for KIAAD\$48 protein partial cds	MT
29088	3.0149440394	F13700	Homo sepiens ribonuclease P protein subunit p40 (RPP40) gene complete cds	other
22960	3.0141662421	T10272	EST:	other
33585	3.0121672451	W93000	EST:	other
220	3.0109180714	D13627	Human mRNA for KIAAD002 gene complete cds	TM
4298	3.0024671064	U36448	Human Ca2+-dependent activator protein for secretion mRNA complete cds	TM
7445	2.9995543641	AA104023	ESTs	,
40903	2.9990347068	N68670	EST:	7
18055	2.9973386648	AA179387	EST ₆	other
7282	2.9952792596	AA083339	ESTA	other

FIGURE 8 (cont.)
23 of 37

9348	2.9949017671	H03686	EST.	TM
806	2 9877 476515	D87009	Human (lambda) DNA for immunoglobilin light chain	7
38447	2.9876031644	AA504255	Human protein kinase ATR mRNA complete cda	other
41454	2.9870604981	R46837	EST:	7
9662	2.9869352306	L19161	TRANSLATIONAL INITIATION FACTOR 2 GAMMA SUBUNIT	other
16978	2.9801154057	AA063625	EST	7
37426	2,9756408909	AA454016	EST:	other
2588	2,9725898298	M27878	Zinc finger protein 84 (HPF2)	other
15174 33620	2.9695024379 2.9657446567	U82987 W93943	Human 8d-2 binding component 3 (bbc3) mRNA partial cds ESTs	other
6784	2.965506112	Y11681	Homo sapiens ribosomel protein S12 gene nuclear gene	7
41077	2.9642389716	N95028	encoding matochondrial protein complete cds ESTs	TM
1932	2.9609985996	L24804	Human (p23) mRNA complete cds	other
39556	2.9588964022	F03738	EST:	other
16106	2.9574232912	AA002258	ESTs .	SS.
32156	2.9574232912	R40381	EST:	7
13617	2.9552305838	AA456646	ESTs	other
11989	2.955203991	AA281251	ESTs Weakly similar to inthorax protein trxlf	other
6056	2.947654132	X58194	[D.melanogaster] Pantophysin (human keretinocyte line HeCaT mRNA 2106 nt]	тм
15446	2.9445456286	W27374	Homo sapiene 10kD protein (8C10) mRNA complete cds	other
38085	2.9445277634	AA482557	EST	7
13878	2.9444133384	AA478604	EST	other
6209	2.9422425032	X76770	H.sapians PAP mRNA	other
388	2.9357591919	D28791	Phosphatidylinositol glycan class A (paroxysmal noctumal hemoglobinuna)	7
1351	2.9266145582	HG4755-HT5203	EST - HG475S-HT5203	7
42624	2.9266145582	W87804	EST*	other
34895	2.9242794509	AA311972	EST4	other
20157	2.9214162976	N23393	ESTs	other
29248	2.9188102156	H52918	ESTS	7
4893 10104	2.9178533564 2.9150324884	U66815 R23855	Human SW/SNF complex 155 KDa subunit (BAF155) mRNA complete cds ESTs	other
15039	2.9147218324	U46116		TM
1605	2.9141775797	L00058	Protein tyrosine phosphatase receptor type gamma polypeptide V-myc avian myelocytomatosis viral oncogene homolog	7
4536	2.907560336	U48705	Receptor protein-tyrosine tunesu EDDR1	,
10173	2.905710598	R56678	ESTs Wealty similar to cell division control protein CDC21 (H.sapiens)	7
26555	2.9056210172	AA279071	ESTs Weakly similar to TOBA11.2 [C.elegans]	other
4401	2.9047655582	U41515	Human deleted in spit hand/split foot 1 (DSS1) mRNA complete cds	other
21009	2.8995011918	N90401	EST4	TM
3602	2.894817322	U01317	HEMOGLOBIN EPSILON CHAIN	?
4833	2.8919254016	U63455	Sulfonylurea receptor (hyperinsulinemta)	7
36200	2.6912301426	AA421164	ESTs	7
26845	2.8898309441	AA281076	ESTs .	other
35299 9504	2.8887661574 2.8880347344	AA398622 M74558	Transcription factor 6-like 1 (mitochondriat transcription factor 1-like) Human Sft. mRNA complete cds	other
5216	2.8877977515	U83410	Human CUL-2 (cut-2) mRNA complete cds	other
12313	2.8847621603	AA397916	ESTs	other
5928	2.8836060438	X62048	WEE1-LIKE PROTEIN KINASE	7
39586	2.6818258313	F09155	ESTs	TM
34758	2.8775214637	AA287680	EST	2
18199	2.8753849024	AA195318	ESTs	other

FIGURE 8 (cont.)
24 of 37

19867	2.8720974689	H61476	EST:	7
1809	2.667937293G	X69398	CD47 entigen (Rh-related entigen integrin-associated signal transducer)	SS.TM
S254	2.862067239	U86782	Human 26S protessame-essociated pad1 homolog (POH1) mRNA complete cds	other
13579	2.6570520494	AA455967	Human neuronal PAS2 (NPAS2) mRNA complete cds	,
1117	2,8568053461	HG3075-HT3236	EST - HG3075-HT3236	7
20533	2.8564678641	H54407	ESTs	TM
38495	2.8562453397	AA505118	Human nucleaporin 98 (NUP98) mRNA complete cds	other
33729	2.8548155651	Z39654	EST	,
2028	2.8532778139	L35035	RIBOSE 5-PHOSPHATE ISOMERASE	other
27374	2.8520674335	AA425816	ESTs Weakly similar to Y53C1ZA3 [C.elegans]	other
19404	2.6516690748	H20568	ESTs	other
26108	2.8504706329	AA243189	ESTs	SS.
4189	2.8439972255	U30930	UDP glycosytransferate 6 (UDP-galactose ceramide galactosytransferase)	TM
16708	2.8427388072	AA043944	ESTs	other
357	2.8350474214	D26156	Human mRNA for transcriptional activator hSNF2b complete cds	other
26045	2.8315740098	AA236276	EST:	other
17796	2.6312342777	AA150435	ESTs	other
6059	2.6288722809	AA310967	ESTs Weakly similar to TO4AB.11 [C.elegans]	other
40914	2 827999584	N69220	EST6	other
27169	2.8263163852	AA410267	H.sapiens mRNA for basic transcription factor 2 34 kD	other
21358	2.8262413945	R16079	subunit ESTs	recto
3572	2.8261469131	\$87759	Protein phosphatase 2C alpha (human teratocarcinoma	Other
11677	2.8259099942	AA262727	mRNA 2346 nt) ESTs	other
1653	2.8234017508	L05424	CD44 entigen (cell adhesion molecule)	7
24645	2.8131264428	239106	ESTs .	ather
35830	2.8126257031	AA411448	EST:	TM
4433	2.6114422177	U43279	EST - U43279	7
20151	2.8109454503	N22895	Home sapiens clone 1400 unknown protein mRNA	other
38648	2.8084431065	AA599267	partial ods EST - RC_AA599267	other
7777	2.8071817929	AA236820	EST:	other
32845	2,80583194	W31566	EST	7
28258	2.8043934162	AA505133	ESTs ·	other
6853	2.798263202	222951	TRANSCRIPTION FACTOR PGS	7
35944	2,7913872996	AA412488	ESTs	7
30648	2.7866523676	N50971	ESTs .	7
18965	2.7857482775	H01411	ESTs .	TM
8616	2.785444221	AA460077	ESTe '	other
14945	2.7838257917	T99606	ESTs Weakly similar to F35G2.2 [C.elegans]	other
8375	2.7805657722	AAA22160	H.sapiens NAP (nucleosome assembly protein) mRNA	other
34929	2.7792111121	AA342084	complete cds EST - RC_AA342084	other
326	2.7766978435	D21262	Human mRNA for KIAA0035 gene partial cds	other
27057	2.7781218063	AA400998	ESTs	SS.
36292	2.7748002184	AA424513	EST - RC_AA424513	other
6480	2.7735431318	X91788	H.sapene mRNA for Icin protein	other
15424	2.7731675808	W27054	APOLIPOPROTEIN AI REGULATORY PROTEIN-1	other
11602	2.7730818255	AA243007	EST:	7
18175	2.77056688	AA194730	EST:	,
25202	2.7698585996	AA034527	EST	7
1681	2.7697545972	L07493	Replication protein A (E coli RecA homolog RAD51	other
14566	2.767984858	AA621122	homolog) ESTs	other
25814	2.7633374335	AA115769	EST.	other

FIGURE 8 (cont.)
25 of 37

141	82 2.3	7506048934	AA490885	ESTs	ather
315	99 2.1	7591187958	N72196	EST	other
182	53 2.7	7471964081	AA206370	ESTs	other
61	23 23	7442487702	×76092	Regulatory factor (trans-acting) 3	other
229	11 2.5	7433449859	T03865	EST ₃	other
355	49 7.	743246906	AA401274	Homo sapiene RRM RNA binding protein Gry-top (GRY-	other
359	55 2.3	7389431758	AA412528	RBP) mRNA complete cas ESTs Weakly similar to ORF2 consensus sequence encoding endonuclease and reverse transcriptese minus	other
176	42 2.7	7377607284	AA132983	RNaseH (R.norvegiczs) ESTs Moderately similar to C-1-TETRAHYDROFOLATE SYNTHASE CYTOPLASMIC [H.sapiens]	cther
61	11 2.3	7371784571	X72841	Human retnoblastoma-binding protein (RbAp45) mRNA complete cds	other
414	29 2.	7347564467	R44994	ESTs	other
170	52 2	7323944161	AA070815	EST - RC_AA070815	other
342	43 2.3	7294147034	AA235050	ESTs	7
729	37 2.1	7284347248	T10065	Homo sepiens TLS-associated protein TASR-2 mRNA	other
51	33 2.7	7243199196	U82130	complete cds Human tumor susceptibility protein (TSG101) mRNA complete cds	other
308	37 2.3	7231409239	N54416	ESTs	other
162	43 2.3	7228028265	AAD12902	EST6	TM
199	54 2.3	7215103495	H80100	ESTs	other
64	u 2.	720441384	X89750	H.sapiens mRNA for TGIF protein	other
59	16 2.3	7192579481	X61072	Human mRNA for T cell receptor clone IGRA17	SS.
62	10 2.1	7168544194	X78627	H.sapiens mRNA for translin	7
421	16 2.7	7144176156	T69924	EST - RC_T69924	other
77)1	107230468	AA215333	ESTs .	TM
175	68 2.3	7096978968	AA128905	ESTs	TM
425			W73189	Homo sapiens protein-tyrosine kinase EPHB2v (EPHB2) mRNA complete cots	
298			N21111	EST• .	other
388		7067394943	AA609458	ESTs	other
103			R88880 S54641	ESTs Moderalely similar to zinc finger protein [M.musculus] HZF-16	other
329			V/38366	Human mRNA for KIAA000S gane complete cds	other
352			AA398507	FSTs	other
101	-		R60100	ESTs	?
					: other
325 345			T27697 AA262768	Human mRNA for KIAA0036 gene complete cds	TM
					other
132			AA443720	ESTs .	other
64			AA443460	ESTs	
77			AA236771	ESTs	other
104			AA007234	ESTs	7
11:			HG3132-H13308 M16837	EST - HG3132-HT3308 Human homeo box c1 protein mRNA complete cds	TM
			M 10837 F 10868		7
109			AA291759	Human SH3 domain-containing protein SH3P18 mRNA complete cds ESTs	r _M
418			T33311	Neuronal pentraxin II	other
20			L33881	Protein kinase C icts	7
336			238501	ESTs Weakly similar to PROSABLE ES PROTEIN	other
13		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	HG884+HT884	EST - HG884-H7884	7
247			240075	ESTs	other
76			440075 AA192484	ESTs Weakly similar to MITOCHONDRIAL INNER	other
307			N52078	MEMBRANE PROTEASE SUBUNIT 2 [S.cerevisioe] Homo sepiens mRNA for KNAA0637 protein complete	other
212	5A 24	3723253055	R09195	ods Homo sapiens mRNA for KIAA0564 protein partial ods	other
405			N29325	ESTs Highly similar to 47 KD PROTEIN [Pseudomonas	other
	*				

FIGURE 8 (cont.) 26 of 37

			chlororephis)	
25285	2.6685455408	AA045083	VITAMIN K-DEPENDENT GAMMA-CARBOXYLASE	other
9296	2.667949532	D82775	ESTs Weakly similar to unknown [S.cerevisiae]	55.
12174	2 6669305328	AA292128	EST:	other
38357	2.6852770538	AA491265	EST	TM
3154	2.661959680G	M83712	Cholinergic receptor nicotinic alpha polypeptide 5	TM
7363	2.655440738	AA093834	ESTs Weakly similar to HYPOTHETICAL 16.8 KD PROTEIN IN SMY2-RPS101 INTERGENIC REGION (S. corevisiza)	other
1923	2.6530372325	L23808	Matrix metalloproteinase 12 (macrophage etastase)	SS,
24906	2.6527048053	Z41840	ESTs	other
34726	2.6495430564	AA287278	EST:	\$5,
30407	2.6495430564	N45983	ESTs	TM
20408	2.6459891347	N48787	ESTs Moderately similar to till ALU SUBFAMILY SC	other
7158	2.6455059455	AA037206	WARNING ENTRY !!!! [H.sapiens] ESTs	TM
26256	2,6445109705	AA253351	ESTs	?
19822	2,8431968212	H58684	EST4	2
12379	2 6428192941	AA399418	Homo sepiens mRNA for JM23 protein complete coding sequence (clone BMAGE 34581 and BMAGE 45355 and	other -
22698	2.6396306055	R89287	LLNLc110(133Q7 (RZPD Berlin)) ESTs	other
24161	2 6394502284	W58015	EST ₆	other
9558	2.6370149706	H81497	ESTs .	TM
18104	2.6358767288	AA188801	EST	other
24882	2.6357248889	Z41563	ESTs	other
40038	2.6347974764	H69485	EST	other
8865	2.6344645492	AB002359	Human mRNA for KIAA0361 gene KIAA0361 protein	other
22148	2.6288326965	R51831	ESTs	other
4527	2 6277060831	U51990 ·	Human hPro18 mRNA complete cds	other
8394	2.6275394634	AA426156	ESTs .	TM
20422	2.6272599716	N49300	ESTs	other
41602	2.6258613824	R67258	ESTs Moderately similar to rhotekin (M.musculus)	other
612	2.6257836682	D63480	Human mRNA for KIAA0146 pene partial ods	TM
4821	2.619521444	U62801	Human protease M mRNA complete cds	SS.TM
16807	2.617722928	AA053296	EST.	other
15288	2.6173997018	W07562	ESTs Moderately similar to rA8 [R.norvegicus]	other
38023	2.6135617291	AA481066	ESTs moderately summer to two (Allice Payers)	other
	2.6120077647	T91715	ESTs Highly similar to HYPOTRETICAL 103.6 KD	TM
23822			PROTEIN IN COX58-PFK26 INTERGENIC REGION [Saccharomyces cerevisiae]	****
10951	2.6116016519	AA126719	ESTS	other
6150	2.6113980879	X74262	RETINOBLASTOMA BINDING PROTEIN P48	other
39336	2.6109987712	C20945	ESTs Highly similar to HYPOTHETICAL 54.2 KD PROTEIN IN CDC12-ORCS INTERGENIC REGION [Saccharomyces cerevisies]	
17793	2.6102158178	AA150242	ESTs Highly similar to modulator recognition factor 2 [H.sapiens]	other
26891	2 6085 107387	AA292659	ESTs	other
2175	2.607468576	L42621	Homo sapiens Ly-9 mRNA complete ode	TM
10642 15026	2.6048724507 2.6031453592	AA040149 U41816	Human Chromosome 16 BAC clone CIT987SK-A- 270G1 Human C-1 mRNA complete cds	other
7699	2.6019047419	AA215299	Homo sapiens chromosome 19 cosmid R30783	other
8543	2.6019047419	X95654	Homo sapiens mRNA for SCP-1 complete cds	other
20036	2.5093684678	N62122	ESTs	other
11308	2.5993311375	AA207114	ESTs	other
4086	2.5966362866	U24704	Human antisecratory factor-1 mRNA complete cds	Other
38615	2.5963996726	AA598938	EST - RC_AA598938	other
11819	2,5961501969	AA258189 ·	ESTs	other

FIGURE 8 (cont.) 27 of 37

3743	2.5957446266	AA454103	ESTs	other
28270	2.5939657529	AA521186	EST4	TM
5587	2.5932338399	X13482	U2 SMALL NUCLEAR RIBONUCLEOPROTEIN A	other
1984	2.5930132063	H59617	ESTs Highly similar to UBIQUITIN-CONJUGATING ENZYME E2-17 KD (Drosophila melanogaster)	ather
10655	2 5925442731	AA040882	ESTs	7
14053	2,5899324577	AA485147	ESTs Highly similar to HYPOTHETICAL 83.6 KD	other
			PROTEIN ROSD3.2 IN CHROMOSOME III (Ceonorhabdilis elegans)	
31574	2.5883094453	N71303	EST	7
7614	2.5870699315	AA187579	ESTs Weakly similar to Yel007c-ep [S.cerev-size]	other
3797	2.5847445397	AA479195	EST	?
7090	2.5845365105	AA009913	Homo saplens diphthamide biosynthesis protein-2 (OPH2) mRNA complete cds	other
1785	2.5841100415	AA156360	ESTS	other
24219	2.5823376094	WG9960	ESTs	other
19070	2.5813645258	H05970	Human clone 23960 mRNA sequence	other
17719	2.5803606155	AA136569	EST	7
38669	2.579196781	AA599694	Human mRNA for KiAA0133 gene complete cds	TM
20983	2.5763957078	N79565	EST:	TM
9158	2.5731638907	D31446	Homo sepiens breakpoint cluster region protein 1 (BCRG1) mRNA complete cds	other
11362	2.5731137778	AA2272E1	EST:	other
8613	2.5723119462	AA459555	Homo saplens mRNA for KIAA0648 protein partial cds	TM
13866	2.5715997844	AA476319	ESTs	6S.
1030	2.5712815907	R86178	Ataxle telangiectasia mutated (includes complementation groups A C and D)	7
22289	2.567916035	R59601	EST	7
1825	2.5673459608	AA206591	EST - RC_AA206591	other
2055	2.5654242568	N55168	ESTs ·	other
3955	2.5645918108	F03605	PUTATIVE 60S RIBOSOMAL PROTEIN	other
27530	2.5631130948	AA435999	EST:	other
1795	2.5608471476	L13434	Human chromasome 3p21.1 gene sequence complete ods	7
14746	2.5603154966	D50354	Human mRNA for KIAAD007 gene partial cds	other
2993	2.5587815672	M64929	Protein phosphatase 2 (formerly 2A) regulatory subunit 8 (PR 52) sipha isoform	other
19191	2.5545260975	H11297	EST:	other
12986	2.5507999853	AA430032	ESTs Moderately similar to PTTG gene product	7
15452	2.5488533884	W27451	{R.norvegicus} Human Cdc5-related protein (PCDC5RP) mRNA	other
18000	2.5465671712	AA171592	complete cds ESTs	other
24196		W67524	Human protein-tyrosine phosphatase (HU-PP-1) mRNA	TM
4265		W92703	partial sequence ESTs	other
26446	2 544106171	AA258796	EST Weakly similar to putative p150 [H.sapiens]	7
30438	2,5368548574	N47204	ESTs Weakly similar to C50F4.12 [C.elegans]	other
36365		AA425893	ESTs Weakly similar to probable CBP3 protein homolog	other
26135		AA243765	[C.elegans] ERTs	other
41885	2.200	T23449	ESTs Moderately similar to ZNF127-Xp [H.sapiens]	SS,
15457		W27560	ESTs	other
27748		AA453158	Human kinesin-fike spindle protein HKSP (HKSP)	other
32315		R69840	mRNA complete ods ESTs Weakly similar to LINE-1 REVERSE	2
			TRANSCRIPTASE HOMOLOG [Nyclicobus coucang)	
25310		AA046745	EST	other
42720		239436 AA428204	ESTs ESTs	other
12939		N52243	EST:	other
30746	2.5198420998 2.5193624578	NS2243 L76703	Home saprens protein phosphatase ZA B56-epsion	omer ?
2722			(PP2A) mRNA complete cds	
11609	2.5191765545	AA243303	ESTs	TM

FIGURE 8 (cont.) 28 of 37

9658	2.5185814336	L 16991	Deoxythymidylate kinese	other
12210	2.5172044581	AA293774	ESTs Weskly similar to PROBABLE TRYPTOPHANYL- TRNA SYNTHETASE MITOCHONDRIAL [C.elegans]	other
3563	2.5169918533	883364	EST - \$83364	other
42407	2.5128230047	W44768	Homo szpiens nephrocystin (NPHP1) mRNA partial cds	7
32826	2 5128052161	W20391	Human mRNA for kinesin-related protein partial cds	other
9693	2.5119977116	L37747	LAMIN B1	?
27862	2.5094571267	AA458908	ESTs	TM
33691	2.509287494	238630	EST	other
17288	2.5088824844	AA085178	EST:	SS,
9888	2.5076170902	N35449	ESTs Highly eimilar to HYPOTHETICAL 30.3 KD PROTEIN IN APE1/LAP4-CWP1 INTERGENIC REGION (Saccharomyces cerevisiae)	other
5932	2.5073880985	X62153	Minichromosome maintenance deficient (S. cerevisiae)	other
15885	2.5053862932	X95073	H, sapiens mRNA for transin associated protein X	çiher
17952	2.5049193223	AA165677	ESTs Weakly smilar to F16A11.1 (C.alegans)	other
12197	2.5042458391	AA293206	EST:	· other
6210	2.5042034458	X76942	Homo sapiens golgin-245 mRNA complete ods	other
34047	2.5041917773	AA194166	ESTs Weakly similar to G1 TO 5 PHASE TRANSITION PROTEIN 1 HOMOLOG [H.saplens]	other
16929	2.5034461307	AA058952	ESTs	other
26834	2.5028075682	AA287138	ESTs Weakly similar to ASPARTYL-TRNA	cther
5157	2.5017270258	UB0034	SYNTHETASE [Thermus equaticus thermophikus] Human mitochondrial intermediate peptidase precursor (MIPEP) mRNA mitochondrial gene encoding mitochondrial protein complete cds	other
38434	2.5005880672	AA497013	ESTs	7
33269	2.5000262771	W72987	EST#	other
26991	2.4990009911	AA398284	ESTe	other
7590	2.4948786183	AA173505	ESTs Weakly similar to HYPOTHETICAL 45.1 KD PROTEIN IN PHO2-POL3 INTERGENIC REGION [S.carevisiae]	other
14960	2.4896232864	U05237	Human fetal Atz-50-reactive clone 1 (FAC1) mRNA complete cds	other
13585	2.4866752902	AA455999	ESTs Highly similar to NEUROLYSIN PRECURSOR [Sus scrots]	other
35901	2.4847673158	AA412151	ESTs	other
38185	2.4826740426	AA487508	Home sapiens mRNA for KIAA0688 protein complete	other
34676	2.4824371274	AA284744	Annexin XI (56kD autoantigen)	other
1424	2.4811113231	J02645	Eukaryotic translation initiation factor 2A	other
16778	2.4800522256	AA047008	EST:	ather
21876	2.4789005203	R43286	EST - RC_R43288	?
17779	2.4695725489	AA149641	EST#	other
24559	2,4682754649	238588	EST&	other
7781	2.467947166	AA242904	Homo sapiens proline-rich Gia protein 1 (PRGP1)	7
7474	2.4677129013	AA126592	mRNA complete cds ESTs Weakly similar to No definition line found	other
34290	2.4675279697	AA236866	(C.elegans) ESTs	other
5316	2.4673813483	U90905	Human clone 23574 mRNA sequence	TM
10216	2.4645666539	R68884	ESTS Highly similar to HYPOTHETICAL 44 2 KD PROTEIN IN SCO2-MRF1 INTERGENIC REGION	other
18109	2.4634292267	AA188981	[Saccharomyces cerevisiae] Homo sapiens retinoblastoma-associated protein HEC mRNA complete ods	7
6485	2.4613518897	X92098	H.sapiens mRNA for transmembrane protein mp24	SS,TM
34954	2.4591845976	AA342959	EST - RC_AA342959	7
42558	2.4588830205	W74751	eY23	other
27444	2,4585750563	AA430160	ESTs Weakly similar to F25H9,7 [C.elegans]	other.
21284	2.4582503599	R10301	EST	,
8920	2.4568596729	AF006265	Homo sapiens cancer associated surface antigen (RCAS1) mRNA complete cds	other
30037	2.4544484116	N27439	ESTs .	TM
27602	2.4527990177	AA443702	ESTs Weakly similar to W02B12.7 [C.elegons]	TM

FIGURE 8 (cont.)
29 of 37

25040 2.4525517032 S59184 RYK receptor-tike tyrosene kinkse 2574	
24487600271	TM
Address	othe
29382 2.4470532391 H72914 ESTs 35521 2.446585249 AA400831 ESTs 20324 2.4464518504 N35406 Phospholopase C bate 4 18630 2.446034893 F02505 ESTs 21067 2.4406971835 R0166 EST 9950 2.4398330157 N71503 ESTs 13665 2.4393228422 N93629 ESTs 15120 2.4345895403 U73524 Human putative ATP/GTP-binding protein (HI mRNA complete cds 15120 2.4345895403 U73524 Human C-1 mRNA complete cds 15120 2.4345895403 U73524 Human C-1 mRNA complete cds 15120 2.4345895403 U73524 Human C-1 mRNA complete cds 15120 2.4328395 AA267115 ESTs 15120 2.42283959 AA267115 ESTs 16073 2.4231729031 AA180453 EST 16073 2.4731729031 AA180453 EST 16073 2.4731729031 AA35698 EST-RC_AA435698 16527 2.41674215 F11087 ESTs 3457 2.4166224787 S74728 Aniquitar 18606 2.4177693475 AA598844 ESTs 20367 2.4157947 N76086 ESTs 24752 2.411498374 Z40012 Homo sapiens mRNA for KIAA0587 protein or document of the complete cds 11701 2.4134093351 AA250301 Homo sapiens RRM RNA binding protein Grym RPJ mRNA complete cds 11672 2.411956031 Z40956 ESTs 24622 2.4119066031 Z40956 ESTs 12672 2.4112720798 AA417067 ESTs 4830 2.4106618618 U63717 Human osteoclast stimulating fector mRNA complete cds 12672 2.4112720798 AA417067 ESTs 4830 2.4063182144 AA610084 ESTs 2574 2.406310553 X77746 Ghatamete receptor metabotropic 3 25872 2.404213441 AA610084 ESTs 25874 2.409310553 X77746 Ghatamete receptor metabotropic 3 25875 2.404213441 AA610084 ESTs 25995 2.404213441 AA610084 ESTs 25995 2.404265999 U43379 Human osteoclast stimulating fector mRNA complete cds 2578 C.AA404995 EST-RC_MAND4995 EST-RC_AA404995 EST-RC_A	oth
2.4465885249	7q21-q31,1 oth
20324 2.4464518504 N35406 Phospholopase C beta 4 18620 2.4460334893 F02506 ESTs 21087 2.4406971835 R00186 EST 9950 2.4396530157 N71503 ESTs 31965 2.4365228422 N93629 ESTs 15120 2.4345895403 U73524 Human putative ATP/GTP-binding protein (HimRNA complete cds 15120 2.4345895403 U73524 Human C in mRNA complete cds 28813 2.439770686 C59257 Human C in mRNA complete cds 33082 2.4295434916 AA482284 ESTs 7960 2.427332589 AA287115 ESTs 18073 2.4231729031 AA180453 EST 18073 2.4231729031 AA180453 EST 18073 2.4187841215 F11087 ESTs 3457 2.4186224787 S74728 Antiquitin 38806 2.4177633475 AA598844 ESTs 20067 2.41519947 N75086 ESTs 24752 2.414198374 Z40012 Homo sapiens mRNA for KIAA0587 protein or cds 24752 2.413994256 AA621611 ESTs 452 2.4135942278 D38076 RAN binding protein 1 11701 2.4134095351 AA253031 Homo sapients RRN RNA binding protein Gryn RBP) mRNA complete cds 24822 2.4119066031 Z40958 ESTs 18622 2.4119066031 Z40958 ESTs 12672 2.4112720798 AA417067 ESTs 4536 2.4106618618 U83717 Human osteoclast stimulating factor mRNA co-cds 24750 2.403828799 T83729 EST - RC_T83729 10987 2.403828799 T83729 EST - RC_T83729 10987 2.407854868 AA132239 EST - RC_T83729 10987 2.4073821434 AA610064 ESTs 10572 2.4073821434 AA610064 ESTs 10585 2.4022664297 AA426291 EST - RC_AA4094 similar to No definition time found for the complete cds 10585 2.402266497 AA426291 EST - RC_AA6094 similar to No definition time found for the complete cds 10586 2.402366599 U49379 Human clacyfglycarol kinase epsilon DGK mR complete cds 10586 2.3998090334 N93514 ESTs 10587 2.3998090334 N93514 ESTs 10588 2.3998090334 N93514 ESTs 10588 2.3998090334 N93514 ESTs 10588 2.39982714 AA011452 ESTs	oth
16820	cthe
2406971835 R00186 EST	\$8.
9950 2.4398530157 N71503 ESTs 31965 2.4363228422 N93629 ESTs 15120 2.4345895403 U73524 Human putative ATP/GTP-binding protein (HimRNA complete cds 28813 2.439770686 C59257 Human C-1 mRNA complete cds 35082 2.42954M916 AA482284 ESTs 35082 2.422869395 AA287115 ESTs 7960 2.427332589 AA287115 ESTs 18073 2.4231729031 AA180433 EST 18073 2.431729031 AA180433 EST 36755 2.4222443392 AA435698 EST - RC_AA435698 18927 2.418741215 F11087 ESTs 3457 2.4188224787 S74728 Antiquitin 38806 2.4177693475 AA598844 ESTs 20967 2.41519947 N76086 ESTs 24752 2.414198374 240012 Homo sapiens mRNA for KIAA0587 protein of cds 452 2.4135942278 D38076 RAN binding protein 1 11701 2.4134095331 AA253031 Homo sapiens RRM RNA binding protein Gry- RBP) mRNA complete cds ESTs Vealdy similar to 265 proteasome subul- (Hisapena) ESTs 4230 2.4106618618 US9717 Octo 2.4053828799 T83729 EST - RC_BA304995 2.4073821434 AA404995 EST - RC_BA304995 2.4073821634 AA402991 EST - RC_BA304995 2.4074873415736 R39349 2.4074873736 R39349	othe
1965 2.4363228422 N39629 ESTs	7
15/20	othe
### RNA complete ods #### C-1 mRNA complete ods ####################################	55.
28813 2,439770696 059257 Human C-1 mRNA complate cds	(HEAB) TM
34723 2.428289395 AA287115 ESTs 7960 2.427332599 AA285277 Homo sapiens brain expressed ring finger procomplete cos 18073 2.4273125931 AA180453 EST 36755 2.4222443392 AA435698 EST - RC_AA435698 18927 2.4187841215 F11087 ESTs 3457 2.4186224787 S74728 Andquitin 38606 2.4177633475 AA598844 ESTs 20967 2.414198374 AM58844 ESTs 24752 2.414198374 A40012 Morno sapiens mRNA for KIAA0587 protein or code 24843 2.413994276 D38076 RAN binding protein 1 11701 2.413694278 D38076 RAN binding protein 1 11701 2.413694278 D38076 RAN binding protein 1 13655 2.412509306 AA458919 ESTs 12672 2.4119206631 Z40958 ESTs 12672 2.4119270798 AA417097 ESTs 42200 2.4083828799 T83729 ESTs Hig	othe
18073 2.427332589 AA285277 Homo sapiens brain expressed ring finger procomplete cas	olte
18073 2.4731729031 AA180453 EST	OU're
18073 2.4231729031 AA180453 EST 36755 2.4222443992 AA435698 EST - RC_AA435698 18927 2.4187841215 F11087 ESTs 3457 2.4186224787 S74728 Antiquitin . 38806 2.4177633475 AA598844 ESTs 20067 2.41519947 N76086 ESTs 24752 2.414188374 240012 Homo sapiens mRNA for KIAA0587 protein of cids 24752 2.414188374 240012 Homo sapiens mRNA for KIAA0587 protein of cids 24843 2.4138974256 AA621611 ESTs 452 2.4135942278 D38076 RAN binding protein 1 11701 2.4134035331 AA253031 Homo sapiens RRM RNA binding protein Gry RBP) mRNA complete cids 13655 2.412509306 AA458919 ESTs Weakly similar to 26S proteasome subulification of the cids o	protein mRNA othe
18927 2.4187841215 F11007 ESTa 3457 2.4186224767 S74728 Andquitin 38606 2.4177633475 AA598844 ESTa 20067 2.41519947 N76086 ESTa 24752 2.414188374 240012 Homo sapiens mRNA for KIAA0587 protein of cidal	othe
3457 2.4186224767 S74726 Antiquitin	othe
38006 2.4177633475 AA598844 EST a 20967 2.41519947 N76096 EST a 20967 2.41519947 N76096 EST a 20967 2.41519947 N76096 EST a 24752 2.414198374 Z40012 Mome asplenis mRNA for KIAA0587 protein of cides 28443 2.4135942278 D38076 RAN binding protein 1 452 2.4135942278 D38076 RAN binding protein 1 11701 2.4134095351 AA253031 Home asplenis RRIM RNA binding protein GryRBP) mRNA complete cids 13655 2.412509306 AA458919 EST in Wealdy similar to 265 proteasome subuly [H. saperus] 24622 2.4119066031 Z40956 EST in Laperus] 24622 2.4119066031 Z40956 EST in Laperus] 42838 2.4106618618 U63717 U63717 42900 2.4083828799 EST - RC_BA720 EST - RC_BA72019 10987 2.4073821434 AA604995 EST - RC_BA740493 2572 2.4073821434 AA604995 EST - RC_AA40493	othe
2-41519947	TM
24752 2.4141498374 240012 Homo sapiens mRNA for KIAA0587 protein of cids 28443 2.4138974256 AA621611 ESTs 452 2.4135942278 D38076 RAN binding protein 1 11701 2.4134095351 AA253031 Homo sapiens RRM RNA binding protein GryRBP) mRNA completis cds 13655 2.412509306 AA458919 ESTs Weakly similar to 26S proteasome subulification 24822 2.4119066031 Z40956 ESTs 12672 2.4112720798 AA417067 ESTs 4838 2.4106618618 U63717 Human osteoclast stimulating factor mRNA colonic material factor minal factor m	cine
28443 2.4138974256 AA621611 ESTs 452 2.4138974256 D38076 RAN binding protein 1 11701 2.4134095351 AA253031 Homo appiers RRM RNA binding protein GryRBP) mRNA complete cds 13655 2.412509306 AA458919 ESTs Wealdy similar to 265 proteasome subulification of the complete cds and the complete cds are cd. Called are cd. Called are cd. Called are cd. Called are c	cine
452 2.4135942278 D38076 RAN binding protein 1 11701 2.4134095351 AA253031 Homo sapiens RRM RNA binding protein Gry-RBP) mRNA complete cds 13655 2.412509306 AA458919 ESTs Vealdy similar to 26S proteasome subusification of the control of the contro	complete other
11701 2.4134095351 AA253031 Homo apjens RRM RNA binding protein Gry RBP) mRNA complete dis 13655 2.412509306 AA458919 ESTs Wealdy similar to 26S proteasome subulification of the control	,
13655 2.412509306 AA458919 ESTs Weakly similar to 26S proteasome subused to 24422 2.4119066031 Z40956 ESTs Weakly similar to 26S proteasome subused to 24422 2.4112720798 AA417097 ESTs ESTs 4838 2.4106618618 U63717 Human osteoclast stimulating factor mRNA concept to 2.4083828799 T83729 EST - RC_T83729 EST - RC_T837	othe
24822 2.4119066031 Z40950 ESTs [N.sapana] ESTs [N.sapana] ESTs 42672 2.411270798 AA417097 ESTs 4830 2.4106618618 U83717 Human osteoclast stimulating factor mRNA co cots 42700 2.4083828799 T83729 EST - RC_T83729 10987 2.4078548068 AA132239 EST - RD_T83729 10987 2.4073821434 AA40499S EST - RD_T83729 EST - RD_T83729 2.4073821434 AA40499S EST - RD_T83729 EST - RD_T83729 2.4073821434 AA40499S EST - RD_T83729 EST - RD_T83729 2.403310553 X77746 GALdamete receptor metabotropic 3 2.403213441 AA610064 ESTs 2.4031905697 AA26291 ESTA Weakly similar to No defindion tine found (C.elegans) 2.4031905697 N93403 ESTs 2.4031905697 AA27745 ESTs 2.3998090334 N89514 ESTE Weakly similar to oxidoreductase (H.sap 2.3986103066 H99972 ESTs 3.50766 2.3971559161 AA435815 Human Clik-associated RS cyclophilin CARS-Complete cds 2.39947415736 N93185 ESTs 7097 2.39382714 AA011452 ESTs	
12672 2.4112720798 AA417097 ESTS 4838 2.4106618618 U83717 Human ostacolast attimutating factor miRNA colors of the color o	•
4836 2.4106618618 U83717 Human osteoclast stimulating fector mRNA co- cote cote cote cote cote cote cote cote	othe
42700 2.4083828799 T63729 EST - RC_T83729 10987 2.4078548688 AA132239 EST - RC_T83729 10987 2.4078548688 AA132239 EST - RC_T83729 10987 2.4078548688 AA132239 EST - RC_T83729 10987 2.4073821434 AA04995 EST - RC_AA04995 10987 2.405310553 X77746 GALAMARIE RESTRICT RE	other
10987 2.4078548688	complete other
PROTEIN IN SHP1-SEC17 INTERGENIC REST	,
35672 2.4073821434 AA04995 EST - RC_AA04095 6224 2.405310553 X77746 Glutamate receptor metabotropic 3 28395 2.404213441 AA610064 ESTs 63690 2.4032664297 AAA26291 EST6 Weakly similar to No defindion time found (C.elegans) 21045 2.4031905697 N93403 ESTs 4558 2.4024665999 U49379 Human diacy/glycarol kinase epailon DGK mR complete dds 12916 2.3998090334 N93514 ESTs Weakly similar to oxidoreductase [H.sap 29769 2.3986103066 H99972 ESTs 63786 2.3971559161 AA435815 Human Cik-associated RS cyclophilin CARS-Complete dds 13942 2.3947415738 N93185 ESTs	.4 KD other REGION
28395 2.40213441 AA610064 ESTs 36390 2.4032664297 AA426291 ESTs Weakly similar to No defindion line foliax [C.elegans] 21045 2.4031905697 N93403 ESTs 4558 2.4024665999 U49379 Human diacytglycarol kinase epsilon DGK mR complete cds 12916 2.3998505067 AA427745 ESTs 20850 2.3988090334 N69514 ESTs Weakly similar to oxidoreductase (H.aap 29769 2.3986103066 H99972 ESTs 31942 2.3947415736 N93185 ESTs 7097 2.39382714 AA011452 ESTs	other
36390 2.4032664297 AAA26291 ESTs Weakly similar to No defindion tine found [C.elegans] 21045 2.4031905697 N93403 ESTs 4558 2.4024665999 U49379 Human diacytglycarol kinase epsilon DGK mR complete cds 12916 2.3998505067 AA427745 ESTs 20850 2.3998090334 N69514 ESTs Weakly similar to oxidoreductase (H.sap 29769 2.3986103066 H99972 ESTs 36786 2.3971559161 AA435615 Human Cik-associated RS cyclophilin CARS-Complete cds 31942 2.3947415736 N93185 ESTs 7097 2.39362714 AA011452 ESTs	TM
C.elegans C.elegans	other
21045 2.4031905697 N93403 ESTs	and other
4558 2.4024665999 U48379 Human diacytglycarol kinase epsilon DGK mR complete dds 12916 2.3998505067 AA427745 EST E Complete dds 20850 2.3988090334 N69514 EST E Wealdy similar to oxidoreductase (H. sap 29769 2.3986103066 H99972 EST E 36786 2.3971559161 AA435615 Human Cik-essociated RS cyclophilin CARS-Complete dds 31942 2.3947415736 N93185 EST s 7097 2.39362714 AA011452 EST s	,
12916 2 3998505067 AA427745 ESTs 20850 2.3988090334 N69514 ESTs Weakly similar to oxidoreductate (H. sap 29759 2.3966103066 H99972 ESTs 36786 2.3971559161 AA435815 Hunna Cik-associated RS cyclophilin CARS-Complete cds 31942 2.3947415736 N93185 ESTs 7097 2.39382714 AA011452 ESTs	
29769 2.3968103066 H99972 ESTs 36786 2.3971559161 AA435615 Human Cik-associated RS cyclophilin CARS-C complete cds 31942 2.3947415736 N93165 ESTs 7097 2.39362714 AA011452 ESTs	other
29769 2.3965103066 H99972 EST e 36786 2.3971559161 AA435615 Human Ciki-essociated RS cyclophilin CARS-Complete cds 31942 2.3947415736 N93185 EST e 7097 2.39362714 AA011452 EST s	apiens] other
31942 2.3947415736 N93185 ESTs 7097 2.39382714 AA011452 ESTs	other
7097 2.39362714 AA011452 ESTs	-Cyp mRNA other
	other
39462 2.3936147708 D60063 ESTs	other
	other
14420 2.3919915706 AA600322 ESTs Highly similar to AAC-RICH MRNA CLO	LONE AAC3 other
PROTEIN (Dictyostelium discoideum) 34629 2.3916035475 AA282527 EST - RC_AA262527	other
27431 2.3905463084 AA429038 ESTs	TM
6387 2.3904071556 X85372 H.sapiens mRNA for Sm protein F	other
11342 2.3502176276 AA223874 Homa sapiens mRNA for KIAAD704 protein pai	partial cds other

FIGURE 8 (cont.) 30 of 37

1497	2,388369765	J04068	Topoisomerase (DNA) II slohe (170kD)	cther
9841	2.3341922016	M95724	Cantromera autoantigen C	other
11454	2.3820201875	AA233854	ESTs	TM
29950	2.3807499489	N24902	Homo sapiens mRNA for E1B-S5kDa-associated protein	TM
8396	2.3807187289	AA426176	ESTs Weakly similar to Similar to S.cerevisian	other
32978	2.3805995259	W42768	hypothetical protein L3111 (H.sapiens) Human terminal transferase mRNA complete ods	Dine/
27672	2.3784145648	AA459254	ESTs	other
11623	2.3769685069	AA243617	EST:	other
26582	2.3766957777	AA279768	ESTs	other
22142	2.3761275381	R\$1382	Homo sapiens mRNA for KIAA0659 protein partial cds	other
13533	2 3759359566	AA454607	ESTs Highly similar to MYPOTHETICAL 40.2 KD PROTEIN K12H4.3 IN CHROMOSOME (II [Cegnorhabdita sispans]	other
11534	2.3747649776	AA236223	EST+	ather
5976	2.3709397882	X64229	DEK PROTEIN	other
6231	2.3680994679	X76121	Chorolderemie	TM
2362	2.3677644584	M16967	Coagulation factor V	other
22887	2,3673034941	T03314	ESTe .	TM
24371	2.3663729415	W87415	ESTs Weakly similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII (H. sapiens)	other
25286	2.3656134948	AA045261	ESTs	other
9054	2.3647542793	C02472	ESTs Weakly similar to EBV-INDUCED G PROTEIN- COUPLED RECEPTOR 1 PRECURSOR [Nomo sabiens]	other
8163	2.3646144577	AA357394	ESTs .	other
12233	2.364077771	AA343513	ESTs Weakly similar to LINE/lig H-chain fusion protein (M.musoutus)	SS.
22924	2.3634007127	T08195	ESTs.	other
14371	2.361524453	AA599219	ESTs Moderately similar to ALR [H.saptens]	other
12401	2.3607293644	AA400229	ESTs	other
26169	2.3599633182	AA251089	ESTs Weakly similar to ORF YOR281c (S.cerevisiae)	?
23065	2.3592943521	T23539	ESTs Highly similar to zinc (inger protein (M.musculus)	other
20524	2.358218239	N53965	ESTs	other
20837	2.3577032218	N69263	ESTs Highly similar to HEXOKINASE TYPE I [Homo	other
18201	2.3573132815	AA195398	sapiens) Home sapiens DNA sequence from PAC 434014 on chromosome 1922.3-41. Contains the HSD1181 gene for hydroxysteroid (11-beta) Dehydrogensee 1 the ADDRAZBP adenosine A2b receptor LIXE pseudogene the IRF6	other
7813	2.3566868562	AA248297	ESTE	TM
21195	2.3547018746	R07210	EST:	other
13377	2.3513919997	AA449720	Homo sapiens clone 24706 mRNA sequence	other
9714	2.3497245732	L44367	EST.	other
41537	2.3460892052	R55673	EST:	other
17352	2.34595172	AA100925	EST.	other
11914	2.3446813991	AA278907	ESTs	7
24890	2.3440589932	241634	ESTs .	other
28795	2.3434458024	D51272	EST - RC_051272_s	7
36798	2.347525534	AA435870	ESTs Weakly similar to 80564.1 (C.elegans)	other
22491	2.3409294581	R70012	EST	other
4798	2 3403776443	U61538	Human calcium-binding protein chp mRNA complete cds	other
40847	2.3397210986	N66354	EST»	Chec
15657	2.3392349306	W63627	Small inducible cytokine A5 (RANTES)	TM
24482	2.3374045148	Z38137	ESTE	other
42022	2,3336939603	T53138	Homo sepiens mRNA for hTCF-4	TM
38233	2.3314220199	AA489023	EST.	other
41221	2.3310635524	R21531	ESTs Weakly similar to Itili ALU SUBFAMILY J WARNING ENTRY IIII (H.sapiens)	other
8053	2.3297250374	AA309880	ESTs	other

FIGURE 8 (cont.)
31 of 37

363	2 3275393529	D26528	Human mRNA for RNA helicase complete ccs	7
26679	2.3241677574	AA281733	ESTs	other
13407	2,3216524472	AA450200	ESTs	TM
17955	2.3160957399	AA166703	EST:	TM
31858	2.3160841803	N90680	EST	,
24092	2.3151511584	W42845	Homo sapiens protein tyrosine phosphatase PIR1 mRNA complete cds	88.
16759	2.3118245547	AA045294	ESTs	other
7851	2,311355404	AA252436	Homo sapiena clone 23797 and 23917 mRNA partial	other
41176	2.3111568749	R09379	ods Natural resistance-associated macrophage protein 2	TM
3860	2.3104335895	U13913	Homolog of Drosophika slowpoke (potessum channel	TM
40886	2.3077403929	N68149	calchum-edivisted) ESTs Weakly striker to EBV-INDUCED G PROTEIN- COUPLED RECEPTOR 1 PRECURSOR [Homo sapiens]	other
19428	2.3068982601	H22949	EST	?
36080	2.3048383557	AA417282	EST - RC_AA417282	other
27264	2,3043527378	AA418389	ESTs	other
13600	2.3031968696	AA456286	ESTs	other
13552	2.3026968375	AA454843	EST:	other
15664	2.3025773291	W67456	ESTs Moderately similar to YY1-associated factor 2	other
26583	2.3025403178	AA279774	(H.sapiens) ESTs	7
37434	2,3013886299	AA454149	EST	?
7833	2.2992574443	AA249300	EST:	other
3674	2.2985613315	U05237	Human fetal Alz-50-reactive clone 1 (FAC1) mRNA	other
33694	2.2984566375	Z38770	complete cas ESTs	other
11178	2.2972286082	AA167436	ESTs	٠,
16977	2.2912855364	AA064616	EST.	other
19709	2.290119924	H57330	EST	7
5946	2.2900738182	X63337	EST - X63337	?
42097	2.2881548729	T66318	Isoleucine-tRNA synthetase	7
24247	2.2681065691	W73010	Ribosomal protein L37	other
40879	2.2870463837	N67816	ESTs Moderately similar to IIII ALU SUBFAMILY SX	other
5875	2.2860441014	X59405	WARNING ENTRY IIII [H.saplens] Membrane cofactor protein (CD46 trophoblast- lymphocyte cross-reactive antigen)	?
22325	2.2850330577	R60777 .	ESTs	other
9621	2.2844572929	J05032	ASPARTYL-TRNA SYNTHETASE	other
9239	2.2823045248	D79100	EST:	other
41997	2.2818672356	T47788	ESTs	other
31105	2.26091752	N63207	EST	7
39565	2.2794194837	F04320	Replication factor C 37-kD subunit	other
7404	2.2793872556	AA094989	Homo sapiens voltage dependent anion channel protein mRNA complete cds	other
6388	2,2768670475	X85373	H.sapiens mRNA for Sm protein G	other
20263	2.2729348551	N31952	ESTs Moderately similar to MYPOTHETICAL 66.5 KD PROTEIN F02A9.5 IN CHROMOSOME III [Caenomabditis elegans]	olher
14520	2 2722894932	AA620307	EST.	other
21197	2 2718358964	R07320	EST:	other
28203	2.2692501412	AA490969	ESTs	other
38320	2.2687130032	AA490611	EST:	other
41625	2,2680307053	R69333	ESTs	other
4674	2,265734645	U54999	Human LGN protein mRNA complete cds	other
28861	2.2637023919	D60037	EST Weakly similar to C50B8.3 (C.elegans)	other
31062	2.2633840539	N62827	ESYs	olher
26756	2.2627797292	AA283832	ESTs	other
11567	2.2614480815	AA236747	Homo sapiens mitogen activated protein kinase	other

FIGURE 8 (cont.) 32 of 37

				TM
25050	2.2605083659	AA011134	ESTs Weakly smiler to renin (H.sapiens) Human serine kinase mRNA complete cds	other
41935	2.2593192037	129681		other
26895	2 2582367069	AA292765 N34891	H.sepiens mRNA for M-phase phosphoprotein mpp5 Homo sepiens mRNA for KIAA0595 protein partial cds	other
40585	2.2581993468		,	
3343	2.2568482074	M97936	SIGNAL TRANSDUCER AND ACTIVATOR OF TRANSCRIPTION 1-ALPHABETA	other
42435	2.2532463427	W46994	EST.	7
5937	2.2489783488	X62534	High-mobility group (nonhistone chromosomal) protein 2	ather
21241	2.2477801609	R08617	ESTs	TM
25756	2.2472586561	AA135868	EST.	TM
34184	2.2459308213	AA227959	Human cysteine protease Mch2 isoform alpha (Mch2) mRNA complete cds	other
6672	2.2450884129	AA477046	ESTs	other
7387	2.2447544716	AA093977	ESTs	other
28822	2.2424116577	D59352	ESTs .	TM
18016	2.2410305445	AA173223	EST.	other
20843	2.239288723	N68352	Homo sepiens mRNA for ATP-dependent RNA helicase	other
10054	2.2387950133	R10266	#46 complete cds ESTS Highly similar to HYPOTHETICAL GTP-BINDING PROTEIN IN PMH0-PAC2 INTERGENIC REGION [Saccharomyces cerevisies]	other
34094	2.2384154308	AA206088	EST.	other
41246	2.2380827238	R27296	ESTs	ather
22634	2.2346537819	R82837	EST:	other
19686	2.2319351858	H48502	ESTs .	\$S,
34568	2.2306030547	AA280609	ESTs Weakly similar to K02B2.3 gene product	other
28448 .	2.2295708871	AA621752	[C.elegans] Human 265 proteasome-associated pad1 homolog (POH1) mRNA complete cds	other
20909	2.2284835116	H71704	ESTs	other
651	2.2260753259	D78129	EST - D78129	SS,TM
40409	2.2244318492	H99877	Homo sepiens exportin t mRNA complete cds	other
20340	2.224062527	N38825	ESTs	other
20002	7.2233023294	H93005	EST - RC_H93005	other
37321	2.2209252793	AA451898	ESTs	other
8274	2.2208752623	AA402095	EST:	other
20221	2.2197714612	N29345	ESTs	other
5792	2.2186801223	X54941	CDC28 protein kinase t	other
4034	2.21806435	U21858	Human transcription initiation factor TFIID subunit TAFII31 mRNA complete cds	other
36222	2.2149577598	AA421481	ESTs .	other
16567	2.2146935655	AA031591	ESTs	ather
4721	2,2134595068	U58046	Human mRNA for KIAA0139 gene complete cds	other
28656	2.2125017907	D19708	Human Gu protein mRNA partial ods	TM
20723	2.2113936194	N66093	E5Ts	other
6714	2,2062571749	Y08612	H sapiens mRNA for Nup68 protein	,
19240	2.205583996	H13265	EST:	other
36447	2.2050784323	AA428168	EST:	other
11688	2.202413216	AA252672	Homo sapiens diphthamide biosynthesis protein-2	other
21650	2.2016153311	R37936	(DPH2) mRNA complete cds Homo sapiene KIAA0440 mRNA partial cds	other
14152	2.2015953698	AA489790	Home sapiens Ran-GTP binding protein mRNA partial	other
42657	2.1975280207	W92771	GLYCINE CLEAVAGE SYSTEM M PROTEIN PRECURSOR	other
4642	2.1968027789	U52427	Human RNA polymerase II subunit hsRPB7 mRNA complete cds	7
32779	2.1962611079	V/02102	EST#	TM
38341	2.1951559134	AA490967	EST:	other
11803	2.1921143838	AA257971	EST:	other
34835	2.190705129	AA292677	ESTs	TM
39085	2.1695804523	AA620599	EST•	other

FIGURE 8 (cont.) 33 of 37

4046	2 1877764122	U22376	MYB PROTO-ONCOGENE PROTEIN	7
11600	2,1876723705	AA242868	ESTs Weakly similar to house-keeping protein	other
5051	2 1866660566	U76638	[M.musculus] Human BRCA1-associated RING domain protein (BARD1) mRNA complete cds	other
33917	2.1864855739	AA167323	EST:	TM
20674	2.1858972155	N63392	EST.	TM
41031	2 1768902734	N91246	EST:	7
25114	2.1759894688	AA020923	EST	7
24711	2.1758363153	Z39645	EST:	other
4733	2.1721786534	U58658	Human unknown protein mRNA within the p53 intron 1 complete cds	other
4871	2 1712198791	U66033	Human glypican-5 (GPC5) mRNA complete cds	other
29733	2.1687028853	H99398	EST	7
23155	2 1678113438	T30550	ESTs ,	other
34638	2.184515923	AA282987	EST	7
35541	2.1621480372	AA400985	Protrymosin slohs	other
1589	2.1598384252	L20591	Annexin III (lipocortin III)	7
15106	2.1591553963	US8111	PROTEIN PHOSPHATASE INHIBITOR 2	7
40131	2.1583553082	H79779	Homo sapiens histone deacetylase 3 (HDAC3) mRNA	other
19516	2.158045763	H29207	complete cds EST	other
4136	2.1577799237	U28014	ICH-2 PROTEASE PRECURSOR	other
20276	2.1548737104	N32919	EST:	other
13292	2,1546709291	AA447621	ESTs Highly similar to 40 KD PROTEIN (Borne disease	other
20666	2.154262609	N63165	virus) ESTe	other
6065	2.1526648242	X68560	Sp3 transcription factor	other
18238	2.1516362853	AA205389	ESTs	other
21627	2 1515999154	R37410	EST	7
3438	2.1502571642	\$72024	Eukaryotic translation initiation factor 5A	?
34648	2.1498935434	AA283772	ACTIVATOR 1 36 KD SUBUNIT	other
5964	2.1488964343	X63657	Follicular lymphoma variant translocation 1	SS.
13250	2 1466085975	AA446459	EST:	other
34370	2.1465845856	AA251829	ESTs Weakly similar to HYPOTHETICAL 27.4 KD PROTEIN IN MER2-CPR7 INTERGENIC REGION	other
27996	2,145312871	AA470156	[Saccheromyces cerevisiae] ESTs Weakly similar to dynein 74K chain cytosolic [R.norvegicus]	\$\$.
4408	2.1398865247	U41745	Human POGF essociated protein mRNA complete cds	ather
4187	2,1395632136	U30868	Human tRNA-guanine transglycosytase mRNA complete	other
10604	2,1366859886	AA069549	cds ESYs	other
34552	2.1340290702	AA279985	Human mRNA for KIAA0372 gene complete cds	other
18360	2.1331897016	AA227119	EST.	other
5223	2.1298428563	U83843	EST - U83843	other
37415	2.1270169134	AA453807	EST	other
14582	2.1260941468	AA621340	ESTs Weakly similar to HYPOTHETICAL 39.6 KD PROTEIN IN MID1-NUP 133 INTERGENIC REGION	other
27756	2.123647107	AA453447	(Saccharomyces cerevisiae) ESTs	other
137 6 7	2.1232866197	AA463745	ESTS Highly similar to PRE-MRNA SPLICING FACTOR	other
5173	2.1232706565	U81554	RNA HELICASE PRP22 (Saccharomyces cerevisiae) Homo sapiens signal recognition particle 72 (SRP72)	other
40029	2.1214337319	H68221	mRNA complete cds Human E2 ubiquitin conjugating enzyme UbcH5B (UBCH5B) mRNA complete cds	other
19972	2.1193721042	н83639	ESTe	other
23301	2.117519655	T52847	ESTs	other
20504	2.1134521605	N52966	EST:	other
40145	2.1132200572	H81391	Human mRNA for histamine N-methyttransferase	other
3461	2.1131164397	\$75256	complete cds EST - 875256	55.
41893	2,1124189285	T23611 ·	ESTs	other

FIGURE 8 (cont.)
34 of 37

39298	2.1092181318	C14805	EST - RC_C14805	other
36021	2 1084566145	AA416876	ESTs Workly similar to TRANSFORMATION- SENSITIVE PROTEIN IEF SSP 3521 [H. capions]	other
6382	2.1077406838	AA424199	ESTs Weekly smilar to C5088.3 (C.elegans)	other
28258	2.1075593303	AA598447	Homo sapiens exportin t mRNA complete ccs	other
5807	2 107 1009331	X55740	5' nucleoudase (CD73)	?
19747	2.106109699	Н53572	ESTs .	other
38155	2 1052335506	AA486777	EST:	TM
924	2.1037724222	HG1112-HT1112	EST - HG1112-HT1112 ,	?
8544	2.1022261514	H72630	ESTs	other
6384	2.1005713227	AA424282	Human 75-kD autoantigen (PM-Sc1) mRNA complete ods	other
25165	2.1005132894	AA027837	Retinitis pigmentose 3 (X-bnked recessive)	SS.TM
24348	2.1000366838	W86469	Tropomyosin alpha chain (skeletal muscle)	7
41401	2.0994968367	R43334	Homo sapiens KIAA0410 mRNA complete cds	other
35340	2.0993762592	AA396900	EST - RC_AA398900	other
10896	2.0990741818	AA112063	ESTs Weakly similar to PRE-MRNA SPLICING HELICASE BRR2 (S.caravisiae)	other
381	2.0974305874	D28473	teoleucine-tRNA synthetase	other
22051	2.0971755	R49047	ESTs Wealdy similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII [H.sepiens]	other
3293	2.096563118	M94893	Tastis specific protein Y-linked	TM
11528	2.0954546212	AA236018	ESTs Weakly similar to unknown (S.cerevisiae)	,
11890	2.0952685865	AA276323	Homo sapiens clone 24606 mRNA sequence	TM
13643	2.0952581265	AA458578	Homo sapiens clone 24477 mRNA sequence	other
19927	2.0952547855	H71829	ESTs	other
36511	2.0927695929	AA429632	ESTs	7
2130	2.0925292202	L40407	Homo sapiens thyroid receptor interactor (TRIP9) gene	?
7193	2.0924678877	AA046768	complete cds Homo sepiens clone TUAS Cri-du-chat region mRNA	TM
5448	2.0921643167	X02751	Neuroblastoma RAS viral (v-ras) oncogene homolog	?
35956	2.0875765163	AA412533	ESTs	curer
7525	2.0870133892	AA149259 ,	ESTs .	other
39592	2.0862891765	F09351	ESTs Weakly similar to weakly similar to S. corvisiae	TM
28029	2.0855738844	AA478479	PTM1 precursor [C.slegans] ESTs	other
18425	2.0855157851	AA232103	ESTs	other
23494	2.0843308862	T70045	ESTs Weakly similar to HYPOTHETICAL 33.2 KD	other
30682	2.0840312831	N56906	PROTEIN F26A3.7 IN CHROMOSOME I [C.elegans]	,
32597	2.0840312831	T47333	Human TFIID subunii TAFII55 (TAFII55) mRNA	r other
			complete cas	
33368	2.0835178514	VVB0814	ESTs	other
10259	2.D829121213	R77527	ESTs .	other
21882	2.0825457608	R43365	ESTs	other
20590	2.0820571859	N58146	ESTs	other
12907	2,0807802388	AA427577	EST:	other
22958	2.0770089467	T10264	EST	other
42044	2.0762746251	158753	ESTs	other
4210	2,0750374179	U31814	Human transcriptional regulator homolog RPD3 mRNA complete cds	other
39	2.074214716	AB003698	Homo sapiens mRNA for Cdc7-related kinase complete cds	other
14350	2.0739236064	AA596831	EST:	TM
29840	2.0729224128	N21680	ESTs	other
25593	2.0715918096	AA113149	Homo sepiens IPL (IPL) mRNA complete cds	other
26071	2 0708411247	AA236860	Protein phosphatase 2A regulatory subunit B' alpha-t	other
26529	2.0699045563	AA276594	EST	7
12154	2.0692192056	AA291293	ESTs	other
18817	2 0684614007	F10077	EST	,

FIGURE 8 (cont.) 35 of 37

6635	2.0674931973	X99585	H sagrens mRNA for SMT3B protein	other
6681	2.066065203	Y00971	Phosphordosyl pyrophosphate synthetase 2	atner
22077	2.0647745388	R49482	ESTs	other
11752	2.0645929355	AA256042	EST.	othe:
41257	2.0634413934	R31680	ESTs	SS.
6904	2.0622381932	234897	Histamine receptor H1	TM
16879	2.050262971	AA056538	ESTs	other
35040	2.0595449295	AA481403	EST:	other
4111	2.0567536207	U26312	Human heterochromatin protein HP1Hs-gamma mRNA complete cds	other
32878	2.0545812272	W37448	ESTs	TM
21743	2.0543668448	R40576	ESTs Moderately similar to IIII ALU SUBFAMILY SX WARNING ENTRY IIII (H. sapiens) ESTs	? other
25968	2.0525018401	AA234935	20.7	
24659	2.0506511898	Z39211	Homo sapiens GDP-L-fucosa pyrophosphorytase (GFPP) mRNA complete ods	other
38030	2.0505994824	AA481148	ESTs	other
61	2.0484705331	AC002115	Cytochrome c oxidase subunit VIb	?
6306 8203	2.0474040935	X81625 AA382517	EUKARYOTIC PEPTIDE CHAIN RELEASE FACTOR SUBUNIT 1 EST - AA382517	? other
34357	2.0469306727	AA251430	ESTs Highly similar to RAS-RELATED PROTEIN RAS-	other
36972	2.0468599712	AA442767	10 (Cents familiaris) Tyrosine 3-monooxygenase/typtophan 5-	other
28156	2.0459278063	AA489057	monooxygenase activation protein bata polypeptide H.sapiens mRNA for nuclear protein SA-2	7
24434	2.045695222	W92787	EST:	other
33508	2.0449481783	W88772	Human DNA sequence from cosmid F0811 on chromosome 6. Contains Dato: BING1 Tapasin RGL2	other
37681	2.0449346104	AA460675	KE2 BING4 BING5 ESTs and CpG Istands H.sapiens mRNA for TRE5	other
27125	2 0448698236	AA405505	Homo sapiens mRNA for putative RNA helicase 3' end	other
3780	2.0445300752	U09851	Zinc finger protein 146 (pHZ-52)	other
9112	2.0443252757	D16611	Coproporphyrinogen oxidase (coproporphyria	TM
6357	2.044244223	AA418921	harderoporphyrla) ESTs Highly similar to RSP5 PROTEIN	other
9133	2.0436113204	D30946	[Saccharomyces cerevisiae] ESTs Highly similar to TRANSLOCON-ASSOCIATED	TM
7519	2.0414123824	AA147425	PROTEIN GAMMA SUBUNIT (Rattus norvegicus) EST - AA147425_4	other
14701	2.0413755305	D59324	ESTa	other
380	2.0411495076	D28423	EST - 028423	7
30571	2.0348528804	N49595	ESTs	other
825	2.0329522889	D87328	Hotocarboxylass synthetase (blotin-[proprionyl-	тм
27744	2.0316041265	AA452818	Coerzyme A-carboxylase (ATP-hydrolysing) ligase) ESTs Weakly similar to HYPOTHETICAL PROTEIN HI0034 (Haemophilus influenzae)	otner
3997	2.0311208335	U19906	Arginine vasopressin receptor 1 (AVPR1)	7
22717	2.0302732387	R91394	EST - RC_R01394	7
377	2.0289078264	D28364	EST - D28364	other
28581	2.0274006652	C21163	. EST	other
11790	2.0269672127	AA256678	ESTs Highly similar to POP2 PROTEIN	other
37931	2.0269058272	AA478523	[Saccharomyces carevisiae] ESTs Weekly similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII [H.sapiens]	other
24678	2.0209818539	239349	ESTs Weakly similar to VACUOLAR ATP SYNTHASE 54 KD SUBUNIT (Saccharomyces cerevisiae)	other
10940	2.0209035614	AA122217	ESTs Weakly similar to HYPOTHETICAL 61.3 KD PROTEIN F25B5.6 IN CHROMOSOME (II (C. elegans)	other
13964	2.0207518872	AA479048	ESTs	7
15665	2.019773566	W67631	Homo sepiens clone 24538 mRNA sequence	TM
28379	2.0189373185	AA509710	EST.	other
13349	2.0172119305	AA449269	ESTs Weakly similar to MITOCHONDRIAL PRECURSOR PROTEINS IMPORT RECEPTOR INdurescore crassal	SS.
7322	2.0167797945	AA090692	ESTs Weakly similar to The KIAA0138 gane product is novel. [H.sapians]	other
29358	2 0165296752	H70641	EST - RC_H70641	3

FIGURE 8 (cont.) 36 of 37

24230	2.016017562	W72276	EST•	other
40212	2.0158778189	H88535	Human clone 121711 defective manner transposon Hamar2 mRNA sequence	?
729	2.01573779	O83778	Human mRNA for KIAA0194 gene panial cds	other
17951	2.0144787235	AA165526	Homo sepiens bifunctional ATP suiturylase/adenosino 5'-phosphosuitate kinase mRNA complete cds	ather
33943	2.0135799277	AA171739	EST	other
5870	2.0118426199	X59244	Zinc finger protein 43 (HTF6)	other
36319	2.0118529739	AA425107	ESTs	ather
25654	2.0097423819	AA126951	ESTs Weakly similar to DNA-directed RNA polymerase [D.melanogaster]	other
16344	2.0090457727	AA018907	EST.	7
8118	2.0090099575	AA328993	ESTs	ctner
29962	2 0087628098	N25228	EST:	TM
32236	2.0078250756	R49327	Natural resistance-associated macrophage protein 2	TM
3279	2.0072427596	M94065	DIHYDROGROTATE DEHYDROGENASE PRECURSOR	TM
16255	2.0065069683	AA013349	ESTs	other
37972	2.0059209236	AA479215	EST - RC_AA479215	TM
41256	2.005858844	R31577	ESTs .	other
34834	2.0050133743	AA292655	ESTa	other
23169	2.0039279023	T33215	EST#	other
29851	2.0034762995	N22145	EST:	other
32862	2	W32519	EST	?

FIGURE 8 (cont.) 37 of 37

		fold upregulated of Tumor		
New Key Number	Accession	over normal	Unigene Descriptor	.]
104660	:AA007160	23	ESTs	_\$S
130016	AA055811	14	transmembrane glycoprotein	SS, TM
104954	AA074514	10	ESTs; Moderately similar to (define not available 4753768) [H.sapiens]	Other
105082	AA143763	7	ESTs; Weakly similar to Similarity to S. Pombe BEM1/BUD5 suppressor [C.elegans]	Other
109141	AA176428	7	ESTs	Other
108893	AA135894	6	retinoic acid induced 3	- TM
108927	AA143493		ESTs; Weakly similar to PLECKSTRIN [H.sapiens]	_Other
109027	AA157818		Human endogenous retroviral protease mRNA; complete cds	Other
133015	AA047036	4	ESTs	Other
114546			ESTs; Moderately similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H.sapiens]	Other
104974	AA056263	***************************************	H.sapiens HUNKI mRNA	Other
108695	AA085918		ESTs	.SS
105049	AA121315 AA132554		ESTs; Moderately similar to mysoin heavy chain 12 [H.sapiens]	· Other
133834	AA147510		Homo sapiens serine protease mRNA; complete cds	Other
109244	AA194237	4	ESTs: Weakly similar to C17H11.6 [C.elegans]	Other
128411	AA007555	3;	ESTs; Weakly similar to transformation-related protein [H.sapiens]	SS, TM
114509	AA043551		ESTs	Other
104888	AA053660	3	ESTs ·	Other
114542	AA055768		ESTs	SS
132718	AA056731		Sjogren syndrome antigen A2 (60kD; ribonucleoprotein autoantigen SS-A/Ro)	Other
104953	AA074157	3	ESTs	Other
132784	AA099589	3	GDP dissociation inhibitor 2	Other
130962	AA102051	3,	transmembrane 4 superfamily member 6	SS, TM
134421	AA122386	3)	collagen; type V; alpha 2	SS
105035	AA128486	3	ESTs	Other
105039	AA130349	3	ESTs	Other
105062	AA134968	3	ESTs	Other
133617	AA148318		Human mRNA for KIAA0059 gene; partial cds	тм

FIGURE 9

			. 44166	ESTs; Highly similar to	
	130335	AA156499		TYPE II-ALPHA REGULATORY CHAIN [H.sapiens]	Other
	105132	AA159501	;	HBV associated factor	Other
	109042	AA159525	3	ESTs	Other
	109043	AA159605		ESTs	Other
	132669	AA188378	3	ESTs; Weakly similar to 60S RIBOSOMAL PROTEIN L22 [H.sapiens]	Other
	135398	AA194075	3	nuclear receptor coactivator 4	Other
ĺ	109344	AA213696	<u>:</u> 3	ESTs	ss
Ì	133221	AA235289	3	ESTs; Highly similar to rap2 gene product [H.sapiens]	_ Other
	114496	AA035611	! : : 2	ESTs; Weakly similar to IIII ALU SUBFAMILY SX WARNING ENTRY IIII [H.sapiens]	Other
	128635	AA043959	;	Itropomyosin 4	Other
	***************************************		<u> </u>	ESTs; Weakly similar to similar to WW/rsp5/WWP domain containing proteins	•-
1	129912	AA047344	2	[C.elegans]	Other
1	104927	-AA058855	2	ESTs	ss
	132821	AA070724	2	CD44 antigen (homing function and Indian blood group system)	Other
	108409	AA075578	2	"zm88h3.s1 Stratagene ovarian cancer (#937219) Homo sapiens cDNA clone IMAGE:54545 3', mRNA sequence"	Other
ľ	133621	AA076138	~ ~~~~	H2A histone family; member Y	Other
	108565	AA085342		ATPase; Ca++ transporting; cardiac muscle; slow twitch 2	TM
ľ	104977	AA088228	2	ESTs	Other
	103777	AA093131		Homo sapiens PAC clone DJ0167F23 from 7p15	- Other
ĺ.	108649	AA112540	2	ESTs	Other
	114692	AA121995	2	ESTs; Weakly similar to Similar to potassium channel protein. (C.elegans)	Other
I.	105063	AA134985	2 }	ESTs	Other
l.	133273	AA147725	2	dendritic cell protein	Other
	128515	AA149044		ESTs; Highly similar to the KIAA0195 gene is expressed ubiquitously. [H.sapiens]	ss
	105182	AA191014		ESTs; Weakly similar to Ydr372cp [S.cerevisiae]	Other
L	109277	AA196332	2	ESTs	Other
ļ.	132608	AA199588		ARP3 (actin-related protein 3; yeast) homolog	Other
 	109380	AA219015	2	ESTs	Other
L	130800	AA223386		ESTs; Weakly similar to katanin p80 subunit [H.sapiens]	Other
	129945	AA232104	2	ESTs; Highly similar to (defline not available 4929579) [H.sapiens]	Other
١.	105305	AA233609		spindle pole body protein	Other

FIGURE 9
(Cont.)

1			
128924	AA234962	2. ESTs	TM
*** * 1 hours		and the second process of the second party of	AND A SECRET SEC.
1		Homo sapiens mRNA fo	r KIAA0887 protein;
114895	AA236177	2 partial cds	Other

FIGURE 9 (Cont.)

FIGURE 10

				-	u
٧		D D	ر		
MINEW Ke	N.Num N.	(が数fold Upregulated of Jumos over normal colon 粉部を	Protect Accession (29)	那m New Key Khinn 光清,经路 food juriegulated of a timos over normal colon 粉漆,这样高的人立在的,这句话,在多年,是一个一个一个一个一个一个一个一个一个一个一个一个一个一个一个一个一个一个一个	NUMBER OF STREET
2451	51	52.6	M21305	Human alpha satellite and satellite 3 junction DNA sequence	2
27030	90	7.4	AA411502	ESTs; Weakly similar to serine protease (H.sapiens)	2
23	2		D13666	Homo sapiens mRNA for osteoblast specific factor 2 (OSF-20s)	SS
25461	19	6.8	AA102520	ESTs; Weakly simitar to heat shock protein hsp4 homolog [H.sapiens]	ž
27665	. 65	6.2	AA453783	ESTs	other
39492	92	6.2	F13673	ESTs	other
28050	Ş	5.6	AA489057	H.sapiens mRNA for nuclear protein SA-2	
31485	85	9.6	N71781	ESTS	other
25606	8	5.4	AA132514	density-regulated protein	other
7000	8	5	274616	collagen; type I; sipha 2	SS
25931	31	4.7	AA236200	ESTS	other
12118	25	4.5	AA291528	ESTs	other
32913	13	4,4	W46810	HMT1 (hnRNP methyltransferase, S. cerevisiae)-like 2	other
26864	25	4.3	AA393804	H beta 58 homotog	other
22514	=	4.2	R79392	ESTS	other
25466	99	4.2	AA112012	lactate dehydrogenase A	TM
32276	76	4.1	R92994	matrix metalloproleinase 12 (mecrophage elastase)	SS
32465	88	4.1	T32108	ESTs	other
22430	8	4	R71082	TFAR19 novel apoptosis-related gena	other
30052	52	7	N32586	ESTs; Weakly similar to Ydr339cp (S.cerevisiae)	other
28354	3	3.9	C14037	ESTs; Weakly similar to Yel7c-ap [S.cerevisiae]	2
29604	8	3.9	H98655	Homo sapiens gene for NBS1; complete cds	¥
27592	92	3.8	AA449417	Homo sapiens mRNA for putative glucosyltransferase; partial cds	2
28691	91	3.8	DS1276	STATHMIN	other
19100	8	3.7	H10933	ESTs	other
5891	9.1	3.5	X60486	H4 histone family; member G	۲
12288	88	3,5	AA398243	ESTs; Highly similar to RSP5 PROTEIN (Saccharomyces cerevisiae)	other
23629	.58	3.5	T68700	ESTs	other
25951	151	3.5	AA236672	ESTs; Weakly similar to DFS7 [H.sapiens]	other
477	1.	3.4	D38583	Human mRNA for calgizzarin; complete cds	٥
11193	93	3.4	AA186897	ESTs	Ψ
27183	63	3.4	AA421562	Homo sapiens secreted cement gland protein XAG-2 homolog (hAG-2R)	SS
32899	66)	3.4	W45728	ESTs; Highly similar to HETEROGENEOUS NUCLEAR RIBONUCLEOPR	other
9576	76	3.3	J03464	collagen; type I; alpha 2	SS
10506	90	3.3	AA027086	ESTs; Highly similar to HYPOTHETICAL 1.4 KD PROTEIN IN UBPS-SPT	other
22064	2	3.3	R51309	ESTs	other
39217	21.	3.3	C21242	calponin 2	other
2613	13	3.2	M29540	CARCINOEMBRYONIC ANTIGEN PRECURSOR	WI
27583	183	3.2	AA449068	ESTs	TW
40031	131	3.2	H83442	calechol-O-methyltransferase	other
10131	31	3,1	R56183	eukaryotic translation initiation factor 3; subunit 6 (48kD)	other
25154	2	3.1	AA043353	ESTs; Highly similar to UBIQUITIN-CONJUGATING ENZYME E2-17 KD 1	other
2000		VO. 10 - 10 - 10 - 10 - 10 - 10 - 10 - 10			

		o o			
ķ	20095	0	2624203	0	ָּע
1	22066	3.6	1676500	COT.	other
?	4,033	2.1	AA40b342	E518	other
ş	10844	9	AA100718	non-specific cross reacting antigen	other
ф Ф	11358	3	AA232104		other
43	17389	c	AA121315	ESTs	other
20	17415	3	AA122386	Collagen, type V, alpha Z'	2
21	23772	9	192735	ESTs	TM
25	25331	9	AA070947	Irapomyosin 4	other
જ	25358	9	AA076138	histone macroH2A1.2	other
ž	27039	9	AA406145	ESTs	SS.TM
22	27261	3	AA425544	Homo sapiens done 23689 mRNA; complete cds	other
98	28785	9	D80946	SFRS protein kinase 1	other
24	32192	3	R67275	coflagen; type XI; slpha 1	other
28	3083	2.9	M77349	transforming growth factor, beta-induced; 68kD	SS
23	5519	2.9	X06700	collagen; type ill; alpha 1 (Ehlers-Danios syndrome type IV; autosomal do	other
9	5562	2.9	X12876	keratin 18	other
-01	11618	2.9	AA251902	Homo sapiens lysophospholipase (LPL1) mRNA; complete cds	other
62	17688	2.9	AA147725	Homo saplens GA17 protein mRNA; complete cds	other
63	18024	2.9	AA188378	ESTs; Highly similar to 6S RIBOSOMAL PROTEIN L22 [Rattus norvegicu	other
ð	20941	2.9	N90933	ESTs	ather
65	13612	2.8	AA458899	ESTs; Highly similar to (defline not available 412715) [H.saptens]	¥
99	17799	2.6	AA157818	Human endogenous retroviral protease mRNA; complete cds	other
29	25344	2.6	AA075182	Stognen syndrome antigen A2 (6kD; ribonucleoprotein autoantigen SS-A/R	2
89	25583	2.8	AA131162	ESTs	other
69	32170	2.8	R61297	eukaryolic translation initiation factor 3; subunit 6 (48kD)	other
9	33586	2.8	238656	coalomer protein complex; subunit alpha	SS
11	2396	2.7	M18728	non-specific cross reacting antigen	other
72	3251	2.7	M93036	membrane component chromosomal 4; surface marker (35kD glycoprotein	other
73	8158	. 2.7	AA372630	Homo sapiens GW112 protein (GW112) mRNA; complete cds	other
74	9207	2.7	D79052	ESTs; Highly stmilar to PROTEIN TRANSPORT PROTEIN SEC61 GAMM	other
75	15051	2.7	U64661	Human poly(A)-binding protein processed pseudogene3	
76	15614	2.7	W63627	ESTs; Moderately similar to IIII ALU SUBFAMILY SP WARNING ENTRY II	ΤM
*	25323	2.7	AA070485	Homo sapiens done 23967 unknown mRNA; partial cds	other
78	25329	2.7	AA070827	ESTs; Weakly similar to KIAA18 (H.sapiens)	ΜL
79	25549	2.7	AA127058	ESTs; Weakly similar to predicted using Genetinder (C.elegans)	¥
80	25584	2.7	AA131165	heterogeneous nuclear ribonucleoprotein A2/B1	other
81	27468	2.7	AA441971	Homo sapiens mRNA for KIAA494 protein; complete cds	ΜT
82	32012	2.7	R31180	ESTs	2
8	38087	2.7	AA488991	Homo sapiens chaperonin containing I-complex polypeptide 1; beta subunit	other
\$	38457	2.7	AA598714	Lon protease-like protein	other
8	39421	2.7	F03974	H.sapiens mRNA for Sop2p-like protein	other
98	3758	2.6	U09587	giycy-IRNA synthetase	other
8	8952	2.6	C00038	ESTS	ΗL
88	12978	2.6	AA431191	ESTs	other

	A	æ			-
	17627	28	AA135804	U POWO CONTRACT CONTR	ָט ווי
	20752	2.6	N68921	FSTe: Weakly similar to peopen in 14 cardans	Σ
	22854	26	Terras	CCT.	omer
	25808	3.6	1999944	2.72	N.
	27169	96.	04448870	5555	other
	28096	9.7	AA4100/8	processome (prosome, macropain) 265 subunit, non-ATPase; 11	other
	20706	0.7	AA430302	ES18, Moderately similar to ubiquitous TPR motif; Y isoform [H.sapiens]	other
	20102	7.0	054288	ESIS	ather
	20030	0.7	239041	ES IS, Weakly similar to DNA-DIRECTED RNA POLYMERASE III LARGE	¥
	37.363	2.6	AA455521	E2F transcription factor 5; p13-binding	other
	39170	2.6	C15324	ESTs	SS, TM
	39251	2.6	D20002	"HUMGS972 Human promyelocyte Homo sapiens cDNA clone pm2344 3".	other
	2767	2.5	M37583	H2A histone family; member Z	other
	5468	2.5	X04347	heterogeneous nuclear ribonucleoprotein A1	other
	9243	2.5	D82348	Homo sapiens mRNA for 5-aminoimidazole-4-carboxamide-1-beta-D-nbon	other
	14791	2.5	T35725	ESTs; Highly similar to HYPOTHETICAL 44.2 KD PROTEIN IN SCO2-MR	other
	14804	2.5	148195	eukaryotic translation initiation factor 3; subunit 3 (gamma; 4kD)	other
	16974	2.5	AA070724	CO44 antigen (homing function and Indian blood group system)	ofher
	20031	2.5	N21085	Homo sapiens androgen receptor associated protein 24 (ARA24) mRNA; c	2
	25484	2.5	AA112679	ESTs: Weakly similar to atternatively spliced product using expn 13A [H.sa	TW
	26830	2.5	AA347359	lysozyme (renal amyloidosis)	SS
1	28068	2.5	AA490212	histone macroH2A1.2	other
	30071	2.5	N33011	replication protein A3 (14kD)	other
	32740	2.5	W31600	von Hippel-Lindau syndrome	other
	35870	2.5	AA416785	heterogeneous nuclear ribonucleoprotein A1	other
-	41908	2.5	T59161	Thymosin; beta 1	MT.
	6011	2.4	X66401	professome (prosome; macropain) subunit; bela type; 9 (large multifunction	2
	9201	2.4	D63079	ESTs: Weakly similar to NADH-UBIQUINONE OXIDOREDUCTASE CHAI	other
١	9218	2.4	D79891	ESTs	TM
	10085	2.4	R32993	ESTs: Highly similar to LEUCYL-TRNA SYNTHETASE; CYTOPLASMIC [other
	10253	2.4	R82411	DEK gene	other
1	11107	2.4	AA159501	ESTs; Moderately similar to RBCK2 (R.norvegicus)	other
İ	11846	2.4	AA262969	ESTs: Weakly similar to similar to Yeast hypothetical protein L8167.12 like	other
	12767	2.4	AA424346	ESTs; Weakly similar to unoporphyrhogen III synthase; UROIIIS [H.sapien	SS
	13772	2.4	AA464708	ESTs; Weakly similar to elternatively spliced product using exon 13A [H.sa	other
	16728	2.4	AA053102	cadherin 17; Li cadherin (liver-intestine)	SS,TM
	17774	2.4	AA156243	ESTs; Highly similar to (defline not available 412715) [H.sapiens]	other
	21386	2.4	R24059	ESTs	other
	25433	2.4	AA099589	GDP dissociation inhibitor 2	TM
	25603	2.4	AA132032	Homo sapiens CAGH1a (CAGH1) mRNA; pertial cds	other
	25791	2.4	AA15980	ELKL motif kinase	other
	26153	2.4	AA252627	ESTs	2
	26852	2.4	AA365527	ESTs; Weakly similar to TLS-associated protein TASR [H.sapiens]	other
	27122	2.4	AA416877	ESTa	other
	28365	2.4	C14090	actin; gamma 1	other

FIGURE 10 (CONT) 4 of 8

)	ı
28628	2.4	025560	Homo sapiens DNA from chromosome 19-cosmid R3879 containing USF2;	other
28687	2.4	DS1241	Homo sapiens mRNA for putative vacuolar proton ATPase membrane sect	SS,TM
29850	2.4	N24968	vacuolar H(+)-ATPase subunit	other
32892	2.4	W45457	ESTs: Moderately similar to neuronal thread protein AD7c.NTP [H.sapiens]	other
33811	2.4	AA173143	heterogeneous nuclear ribonucleoprotein G	other
40121	2.4	H93492	ESTs; Highly similar to villin [H.sapiens]	other
40141	2.4	H94877	ESTs; Moderately similar to putative G-binding protein [H.sapiens]	oger
40167	2.4	H96237	collagen; type XI; alpha 1	other
446	2.3	D38073	minichromosome maintenance deficient (S. cerevisiae) 3	oper
3530	2.3	\$81914	DIFFERENTIATION-DEPENDENT GENE 2	ge
7835	2.3	AA252436	Homo sapiens hysophospholipase (LPL1) mRNA; complete cds	ago
10898	2.3	AA121879	proteasome (prosome: macropain) subunit: beta type: 9 (large multifunction	a dio
10965	2.3	AA134138	ESTs, Highly similar to CYTOSOL AMINOPEPTIDASE (Bos taurus)	other
11015	2.3	AA143763	ESTs, Weakly similar to Similarity to S. Pombe BEM1/BUDS suppressor (C.	other
11895	2.3	AA279420	ESTs. Highly similar to (defline not available 433735) (H.sapiens)	M.
13386	2.3	AA451678	ESTS	other
15464	2.3	W28391	proliferation-associated 2G4: 38kD	other .
17619	2.3	AA135406	ESTs	i di
18225	2.3	AA213696	ESTS	a la
20450	2.3	NS3927	ESTs: Weakly similar to phenylativiamine binding protein (H. sapiens)	,
25308	2.3	AA065227	ESTs: Weakly similar to coded for by C. elegans cDNA vk1c1.3 IC elegans	P P
26590	. 2.3	AA282151	ESTs; Weakly similar to predicted using Genefinder (C.elegans)	o Page
27624	2.3	AA452112	Homo sapiens mRNA for putative thioredoxin-like protein	other
7792	2.3	AA460359	ESTs, Weakly similar to DNA-DIRECTED RNA POLYMERASE II 14 KD P	2
28231	2.3	AA600153	DEKgene	other
28722	2.3	D59711	ESTS	other
30363	2.3	N47956	eukaryotic translation Indiation factor 3; subunit 3 (gamma; 4kD)	other
2928	2.3	W47620	ESTs; Weakly similar to reverse transcriptase related protein (H.sapiens)	other
39585	2.3	H11320	Homo sapiens HRIHFB2115 mRNA; partial cds	¥.
40175	2.3	H3666\$	peptidylprolyl isomerase B (cyclophilin B)	other
40366	2.3	N26691	ESTs; Highly stm8ar to (defline not available 467914) [H.sapiens]	other
40733	2.3		ESTs; Weakly similar to 25 kDa trypsin Inhibitor (H.sapiens)	other
4918	2.2	U68105	poly(A)-binding protein-like 1	۰
5165	2.2	U81607	GRAVIN	other
12242	2.2	AA372018	ESTs	other
3154 .	2.2	AA442768	Homo sapiens inner mitochondrial membrane transfocase TIm23 (TIM23)	2
14276	2.2	AA598450	EST\$	other
15721	2.2	W95348	ESTs	¥
20588	2.2	N62945	Homo sapiens hMmTRA1b mRNA; complete cds	Τ
4021	2.2	W42857	ESTS	other
24250	2.2	W84712	calumenin	other
25245	2.2	AA055768	ESTs	SS
25430	2.2	AA099429	SPLICING FACTOR U2AF 35 KD SUBUNIT	MI
25562	9.9	A 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4		

FIGURE 10 (CONT) 5 of 8

cacherin 17: U cacherin (iver-intestine)
NEUTROPHIL GELATINASE-ASSOCIATED LIPOCALIN PRECURSOR
ESTs. Highly similar to KERATIN: TYPE I CYTOSKELETAL 14 [Homo sa Human mRNA for KIAA89 gene, sartial cds
INTERFERON-ALPHA INDUCED 11: \$ KD PROTEIN
ESTS: Moderatery similar to IIII ALU SUBFAMILT' J VVARINIG ENTRY IIII
ESTS: Wodersky similar to Similarity to Yeast hypotherical protein YOR3160
pigment epithelium-derived factor ESTs. Weakly similar to uroporphymogen III symbass; UROIIIS [H.saplen Hone sapiens done 628 unknown mRNA; complete sequence ESTs. Weakly simitar to deducad smino acid sequence is highly homologo ESTs. Highly simitar to nuclear pore complex protein NUP17 [R.nonveglous eukaryotic translation iatilation factor 3, subunit 7 (zeta; 66167AD)
EST s; Highly similar to (define not avaitable 4454524) [H saplena]
ESTs; Moderately similar to gangitoside-induced differentiation associated 2F transcription factor 5, p13-binding ukaryotic translation initiation factor 3; subunit 6 (48kD) ansferin receptor (99; CD71) Homo saplens mRNA for KIAA746 protein; partial cds AA024462 AA148318 AA161292 AA236972 AA260865 AA111889 H18947 N26259 N91492 N98464
AA412452
AA488433
AA521256
AA621604
D59570
F04674
R48608
W90444
W96151 AA608751 057317 H73484 H76323 N27198 T26799 L09604

1			3	3	
١	1		,		ار
9844	-	2	N33807	ESTs: Highly similar to NEDD-4 PROTEIN (Homo sapiens)	other
14032	32	2	AA486092	ESTs: Weakly similar to CH-TOG PROTEIN [H.sapiens]	M
16395	35	2	AA025673	ESTs: Moderately similar to (define not available 416878) [H.sapiens]	M
17327	27	2	AA112540	ESTs	Σ
23083	83	2	T30881	ublquitin-conjugating enzyme E2E 1 (homologous to yeast UBC4/5)	other
526	25625	2	AA133969	ESTs; Weakly similar to neuronal thread protein AO7c-NTP [H.sapiens]	other
280	28073	2	AA490494	ES1s	other
287	28700	2	D53139	ribosomal protein S28	other
5	29095	2	H27188	collagen-binding protein 2 (colligen 2)	other
32	32191	2	R67083	calnexin	SS.TM
2	32897	2	W45664	5' nucleolidase (CD73)	other
۲	10782	1,6	AA074880	ESTs; Weakly similar to HYPOTHETICAL 85. KD PROTEIN IN CPA2-ATP	other
27	27795	1.6	AA460454	ESTs; Weakly similar to KIAAS12 protein [H.sapiens]	other
38	28706	1.6	D54298	Human mRNA for KIAA255 gene; complete cds	M
క	36414	1.6	AA430186	ESTs	offer
6	9379	1.5	N91067	EST8: Weakly similar to F5SA12.9 (C.elegans)	other
6	7866	1.5	N95507	ESTs; Weakly similar to KIAA319 [H. sapiens]	TM
ř	10656	1.5	AA047290	ESTs	other
ř	14977	1.5	U37546	apoptosis Inhibitor 1	MT
2	27065	1.5	AAA 10294	Human mRNA for KIAA336 gene; complete cds	othe
72	29278	1.5	H72948	biglycan	SS
9	31917	1.5	N96238	ESTs	other
æ	38272	1,5	AA496533	EST8	other
7	41396	1.5	R55342	CD68 antigen	other
4	41966	1.5	T67710	ESTs	2
	6439	1.4	AA436304	Homo sapiens androgen receptor associated protein 24 (ARA24) mRNA; c.	other
7	10311	P'1	AA001936	ESTs	other
۲	10859	1.4	AA112149	ESTs	other
=	11279	1,4	AA213410	ESTs	SS
4	13548	1.4	AA456033	ESTs; Highly similar to HYPOTHETICAL 1.4 KD PROTEIN IN UBPS-SPT	other
•	14340	1.4	AA599653	Homo sapiens TCFL5 mRNA for transcription factor-like 5; complete ods	other
m	32180	1.4	R63727	ESTs	other
3	35187	1.4	AA398722	EST8	other
3	7254	1.4	AA453483	EST8	M
	1344	1.3	HG4757-HT5207	"Oncogene MI-Al4, Fusion Activated"	other
"	5397	1.3	V01516	KERATIN; TYPE II CYTOSKELETAL 6D	SS.TM
-	16272	1.3	AA018922	core promoter element binding protein	other
2	23427	1.3	T70358	ESTs; Highly similar to POL POLYPROTEIN (Simian sarcoma virus)	other
2	25716	1.3	AA150741	gene with multiple spiles variants near HD locus on 4p16.3	other
۳	33499	1.3	W93403	EST3	other
3	36574	1.3	AA434454	ESTs; Weakly similar to ort; hypothetical protein (E.ooii)	SS
*	41548	1.3	R62846	EST8	other
"	2386	1.2	M17863	Tracelly file money design of comments and	
			200	מוספווויונים אוספים ופספים ל (פספוופיסווים)	Ξ

12 C C 12 AA411473 13 AA411473 14 AA417581 15 AA417588 17 AA415888 18 AA415888 19 AA415888 10 AA415888 11 AA41588 12 AA41588 13 AA41588 14 AA41588 15 AA41588 16 AA41588 17 AA41588 18 AA41588 19 AA41588 10 AA41588 11 AA41588 11 AA41588 12 AA41588 13 AA41588 14 AA41588 15 AA41588 16 AA41588 17 AA41588 18 AA41588 19 AA41588 10 AA41588 11 AA41588	<u>u</u>	other	olher	olher	6	other		other	other	other					0		ľ	MI		other	other	other	other	other	SS C WARNING ENTRY IIII [H.sa other		other	other					OF MICHOGO CAN	A ontain tinges 2		omer	ľ					-	other
24427	0	ESTs	adducin 1 (alpha)	ESTs	ESTs	ESTs	ESTs. Weakly similar to putative p15 (H.sapiens)	ESTs	ESTs	ESTs	Homo sapiens mRNA for neuropsin; complete cds	SERUM AMYLOID A4 PROTEIN PRECURSOR	ВРГР	H.sapiens DNA for cyp related pseudogen	ESTs	cerebroside (3'-phosphoadenyhisullale:ga	ESTs	ESTs	Homo sapiens BirnEL mRNA; complete cds	EST	ESTs	ESTs	ESTs	ESTs	ESTs; Moderately similar to IIII ALU CLAS	ESTs; Highly similar to LEUKOTOXIN SE	EST	ESTs	"Homo sapiens plasminogen-like protein (F	Accession not listed in Genbank	SEKINE/THREONINE-PROTEIN KINASE	Homo english and MA Kee MA AND	FOTE Highly similar to MITOCHONDBIA	mitwen-adivated ordein kinase-adivated	FATe	FSTe	7-m13-3 s1 Stratagene nancreas (#93728)	ESTS: Weakly similar to IIII ALU SUBFAMI	ESTs: Weakly similar to neuronal thread pr	ESTs; Moderately similar to unknown [H. sapiens	ESTs	The state of the s	<u> </u>
24427 24427 27089 23177 34622 35325 35325 35325 35325 35325 35325 35325 35325 35325 35325 35325 35325 35325 35325 35325 35325 35325 37523 37523 37523 37523 37523 37523 37523 37523 37523 37525 37523 37525 37523	0	238208	AA411473	W73195	AA347691	AA400273	AA435668	AA598939	R02547	151150	AA401047	M81349	S83198	82905X	N66289	AA004502	H13649	N74690	N78784	AA400013	AA461499	AA479896	AA521342	AA609018	N29963	N40559	R41772	R48580	M93143	X33065	Abedea	W04960	WZ6847	WS8725	AA233898	R41999	AA065081	H40486	AA302772	AA342526	AA400521		AA46449
	8	1.2	1.2	1.2	1.2	1.2	1.2	1.2	1.2	1.2	1:1	1.1	1.1	1.1	1.1	1.1	1.1	1.1	1,1	1.1	1,1	1.1	1,1	1,1	1.1	1,1	1.1	1.1	-			-						•	-	1		•	_
	4	24427	27089	33177	34852	35325	36609	38477	40975	41874	8235	9772	14758	15831	20656	24891	29045	31584	31640	35293	37583	37852	38397	38652	40397	40488	41231	41333	3255	2/42	16222	15282	15363	15606	18435	21736	25306	29111	34743	34801	35355	20070	24505

	Ш	other	olher	other	other	other	SS	other	other	other	other	other	SS, TM	other		L	L		other				SS,TM
	O	ESTs	ESTs	ESTs	Human mRNA for KIAA187 gene; complete cds	ESTs	immunoglobulin superfamily; member 3	ESTs	ESTs	ESTs	ESTs	ESTs	Human clone 23732 mRNA; partial cds	EST	ESTs	ESTs; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens]	fibringen-like 1	ESTs	ESTs	ESTs	ESTs	homogentisale 1;2-dioxygenase (homogentisate oxidase)	librinogen; 8 beta polypeptide
66.	၁						ı	R45512	R49459	T40827	045719	H60824	T89122	W02129	AA401409	AA599209	D14446	AA496980	AA056210	N87590	N54429	R08615	T71012
	8	0.9	6.0	6.0	6.0	6.0	6.0	6.0	6.0	6.0	6:0	6:0	6.0	6.0	6.0	6.0	0.8	0.0	0.8	0.7	0.7	0.7	0.6
	¥	9982	10084	11701	12088	15267	15901	21906	22002	23136	28673	29159	32610	32674	35426	38504	244	14204	25250	9371	20461	41029	41985
		308	310	31.1	312	313	314	315	316	317	318	319	320	321	322	323	324	325	326	327	328	329	83

FIGURE 11 1 of 10

			10,10	,
PRIMEKEY	Fold Upregulated in Turnors	Exemplar Accession	Complete Title	ORF Structuaral
	***************************************		small inducible cytokine subfamily B (Cys-X-Cys); member 5	
134804	12.3	L37036	(epithelial-derived neutrophil-activating peptide 78)	TM
130617	11.4	M90516	glutamine-fructose-6-phosphate transaminase 1	TM
404000	40.0		small inducible cytokine subfamily A (Cys-Cys); member 18; pulmonary and	SS
104209	10.6	AB000221	activation-regulated	- TM
109991	10	H09813	ESTs	
124315	8.3	. H94892	v-ral similan leukemia viral oncogene homolog A (ras related)	Other
132977	8.2	U28686	RNA binding motif protein 3	Other
130407	7.4	N29888	ESTs	Other
116176	6.6	AA463725	ESTs; Weakly similar to KIAA0226 [H.sapiens]	Other
119271	6.5	T16387	ESTs	Other
134711	6.5	X04011	cytochrome b-245; beta polypeptide (chronic granulomatous disease)	TM
134326	6.1	U16306	"Human chondroitin sulfate proteoglycan versican V splice-variant precursor peptide mRNA, complete cds"	Other
125852	5.9	H09290	ESTs; Weakly similar to unknown [H.sapiens]	SS, TM
112169	5.9	R48589	ESTs	TM
132528	5.9	AA283006	chromosome-associated polypeptide C	Other
404000			phosphoribosylglycinamide formyltransferase; phosphoribosylglycinamide	0.5 1
134367	5.7	X54199	synthetase; phosphoribosylaminolmidazole synthetase	Other
111929	5.2	R40057	prominin (mouse)-like 1	SS, TM
102165	5	U18321	Death associated protein 3	Other
130441	5	U35835	protein kinase; DNA-activated; catalytic polypeptide	Other
133282	. 5	U52960	SRB7 (suppressor of RNA polymerase B; yeast) hornolog	Other
100783	4.9	HG3748-HT4 018	"Basic Transcription Factor, 44 Kda Subunit"	Other
122223	4.9	AA436158	ESTs	Other
104660	4.8	AA007160	ESTs	SS
113702	4.8	T97307	ESTs; Moderately similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H.saplens]	Other
101185	4.7	L19872	aryl hydrocarbon receptor	Other
105308	4.7	AA233744	ESTs	Other
107168	4.7	AA621540	ESTs	Other
101201	4.6	L22524	matrix metalloproteinase 7 (matrilysin; uterine)	SS
101809	4.5	M86849	"Homo sapiens connexin 26 (GJB2) mRNA, complete cds"	TM
101478	4.5	M23379	RAS p21 protein activator (GTPase activating protein) 1	Other
104695	4.5	AA012953	ESTs	Other
100365	4.4	D78611	mesoderm specific transcript (mouse) homolog	TM
126819	4.3	AA305538	*EST176522 Colon carcinoma (Caco-2) cell line II Homo sapiens cDNA 5' (end, mRNA sequence."	Other
101880	4.3	M97925	;defensin; alpha 5; Paneth cell-specific	SS
126838	4.2	AA858097	pigment epithelium-derived factor	Other
101684	4.2	M63256	cerebellar degeneration-related protein (62kD)	Other
44700		NOCACA	ESTs; Moderately similar to IIII ALU SUBFAMILY SQ WARNING ENTRY	
117634	4.2	N36421	IIII [H.sapiens]	Other
132109	4.1	AA599801	ESTs	Other
115054	4.1		ESTs	Other
106553	4	AA454967	ESTs	Other
119717	3.9	W69134	ESTs	Other

FIGURE 11 (CONT) 2 of 10

PRIMEKEY	Fold Upregulated in Tumors	Exemplar Accession	Complete Title	ORF Structuarel
131945	3.9	M87339	replication factor C (activator 1) 4 (37kD)	Other
128790	3.9	AA291725	secreted frizzied-related protein 4	SS
120562	3.8	AA280036	ESTs; Weakly similar to W01A6.c [C.elegans]	SS
109517	3.7	D60799	ESTs	Other
102618	. 3.7	U65932	extracellular matrix protein 1	SS
106286		AA434441	frizzled (Drosophila) homolog 7	Other
133640	3.6	D83004	ubiquitin-conjugating enzyme E2N (homologous to yeast UBC13)	Other
100335	3.6	D63391	platelet-activating factor acetylhydrolase; isoform lb; gamma subunit (29kD)	Other
104720	3.6	AA018441	ESTs	Other
		U43701	nbosomal protein L23a	Other
107348	3.6	043701	ESTs; Weakly similar to IIII ALU CLASS A WARNING ENTRY IIII	
134989	3.6	AA236324	[H.sapiens]	. <u>ss</u>
111345		N89820	ESTs	Other
			ESTs; Weakly similar to Similar to NADH-cytochrome B5 reductase	
107053	3.6	AA600147	[C.elegans]	Other
107240	3.5	D59368	ESTs	Other
107129	3.5	AA620553	flap structure-specific endonuclease 1	Other
134846	3.5	AA431505	Homo sapiens mRNA for putative Sqv-7-like protein; partial	TM .
101897	, 3.5	\$58544	sperm associated antigen 1	Other
107151	3.5	AA621169	ESTs	Other
106012	3.4	AA411621	ESTs	Other
101950	3.4	S79219	propionyl Coenzyme A carboxylase; alpha polypeptide	Other
116844		H64938	ESTs	Other
128025		AA937173	ESTs	TM
			leukemla associated gene 1; candidate tumor suppressor frequently deleted	
106785	: 3.4	AA478587	In B-cell chronic lymphocytic leukemia (B-CLL)	Other
104518	3.4	. R09815	proteasome (prosome; macropain) 26S subunit; ATPase; 5	Other
117667	3.4	N39214	ser-Thr protein kinase related to the myotonic dystrophy protein kinase	Other
104954	3.3	AA074514	ESTs; Moderately similar to (defline not available 4753768) [H.sapiens]	Other
105372	3.3	AA236481	ESTs	Other
104896		AA054228	ESTs	Other
113485		T87863	ESTs	Other
127003		AA550806	ESTs; Weakly similar to (defline not available 3882151) [H.sapiens]	TM
100305		D50487	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 8 (RNA helicase)	SS
		1	ESTs; Weakly similar to !!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!	
134722	3.2	: W47183	[H.saptens]	Other
126801	3.2	AA512902	ESTs	SS, TM
133503	3.2	M33195	Fc fragment of IgE; high affinity I; receptor for, gamma polypeptide	SS, TM
132183	3.2	L19183	Human MAC30 mRNA; 3' end	TM
105298	3.2	AA233459	ESTs	TM
		N55514	ESTs; Moderately similar to IIII ALU SUBFAMILY SP WARNING ENTRY IIII [H.sapiens]	TM
111046			ESTs	SS
135309		D25984	committee the committee of the committee of the participation of the committee of the commi	Other
102808	3.2	U90426	nuclear RNA helicase; DECD variant of DEAD box family	
100552	2: 3.1	HG2167-HT2 237	"Protein Kinase Ht31, Camp-Dependent"	Other
,0000		AA804487	ESTs	Other

FIGURE 11 (CONT) 3 of 10

PRIMEKEY	Fold Upregulated in Tumors	Exemplar Accession	Complete Title	ORF Structuars
			ESTs; Moderately similar to coded for by C. elegans cDNA CEESD64F	
116127	3.1	AA459703	[C.elegans]	TM
131904	3.1	AA143019	ESTs; Highly similar to surface 4 Integral membrane protein [H.sapiens]	TM
126547	3.1	U47732	transmembrane 4 superfamily member 3	Other
102823	3.1	U90914	carboxypeptidase D	SS
			Homo saplens cytokine receptor related protein 4 (CYTOR4) mRNA;	Other
132298	3.1	N41849	complete cds	Other
127445	3.1	AA906286	ESTs	SS, TM
134395	3.1	L09717	lysosomal-associated membrane protein 2	Other
105743	3.1	AA293300	ESTs; Weakly similar to semaphorin C [M.musculus]	Other
125827	3.1	A1471525	'YY1 transcription factor	
101229	3.1	L27943	cytidine deaminase	Other
102306		U33317	defensin; alpha 6; Paneth cell-specific	SS
407040	•	774445	"yc82f8.r1 Soares infant brain 1NIB Homo saplens cDNA clone IMAGE:2242	Other
107318		T74445	5, mRNA sequence*	Other
120983	en en en en en	AA398209	EST	TM
134700	. 3	AA481414	golgi SNAP receptor complex member 1	SS
131216	3	D31058	ESTS	9.0
127565	3	D79516	"HUM272B04B Human aorta polyA+ (TFujiwara) Homo sapiens cDNA clone GEN-272B04 5', mRNA sequence."	Other ·
106882	2.9	AA489009	ESTs	Other
130874	2.9	T08287	ESTs	Other
101275	2.9	L37936	:Ts translation elongation factor; mitochondrial	Other
111179	2.9	N67239	ESTs	Other
			"yr73g01.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone	
126086	2.9	H70975	IMAGE:210960 5', mRNA sequence."	Other
105365	2.9	AA236275	ESTs	Other
118789	2.9	N75416	ESTs	Other
115117	2.9	AA256492	H.sapiens PAP mRNA	Other
		HG1103-HT1	"Guanine Nucleotide-Binding Protein Ral, Ras-Oncogene Related"	Other
100484	2.9	103 L40391	Homo sapiens (clone s153) mRNA fragment	Other
		·	The state of the s	Other
103075	2.9 2.8	X59543	nibonucleotide reductase M1 polypeptide procollagen-tysine; 2-oxoglutarate 5-dioxygenase (tysine hydroxylase) 2	Other
132164	·	U84573	 	Other
106716	2.8	AA464962	ESTs	Other
134098	2.8	X06323	Human MRL3 mRNA for ribosomal protein L3 homologue (MRL3	SS
134485	2.8	X82153	cathepsin K (pycnodysostosis)	Other
129634	2.8		ESTS	Other
112207	2.8	R49602	!ESTs	Other
124904	2.8	R86970	ESTS	SS
125471	2.8	AA477571	UDP-glucose ceramide glucosyltransferase	SS
132180	2.8	AA405569	fibroblast activation protein; alpha	. 55 TM
104454	2.8	M84443	-galactokinase 2	Other
134282	2.8	T25508	ESTs	tner
101558	. 2.8	M32011	neutrophil cytosolic factor 2 (65kD; chronic granulomatous disease; autosomal 2)	Other
	2.8	AA173238	:small inducible cytokine A5 (RANTES)	Other
130529	2.0	~~ 173£30	Control to the control of the contro	Other

FIGURE 11 (CONT) 4 of 10

PRIMEKEY	Fold Upregulated in Tumors	Exemplar Accession	Complete Title	ORF Structuaral
109565	2.7	F01930	ESTs	Other
112427	2.7	R62604	ESTs	Other
115188	2.7	AA261819	ESTs	Other
121831		AA425374	ESTs	Other
132454	2.7	W78726	ESTs; Highly similar to serine/threonine kinase [H.sapiens]	Other
116399	4	AA599729	Homo sapiens homeobox protein A10 (HOXA10) gene; complete cds	Other
100409	:	D86957	Human mRNA for KIAA0202 gene; partial cds	Other
		R45698	ESTs	Other
130987		D51095	ESTs	SS, TM
107217		AA748483	Homo sapiens basic transcription factor 2 p44 (btf2p44) gene; partial cds; neuronal apoptosis inhibitory protein (nalp) and survival motor neuron protein (smn) genes; complete cds	SS
118722	2.6	N73563	ESTs	SS
		R11604	"y/47c1.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone !MAGE:132 5", mRNA sequence"	Other
104521	· · · · · ·	AA223386	ESTs; Weakly similar to katanin p80 subunit [H.sapiens]	Other
130800			protein tyrosine phosphatase type IVA; member 2	TM
134415		AA329274 AA621557	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WARNING ENTRY	Other
116461	2.6	HG4297-HT4	IIII [H.sapiens]	
100864	2.6	567	Transcriptional Coactivator Pc4 *zi43h5.r1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone	Other
103818	2.6	AA150614	MAGE:54729 5', mRNA sequence"	Other
105713		AA291321	ESTs	Other
114969		AA250775	ESTs	Other
130415	•	X07290	Human HF.12 gene mRNA	Other
101791	2.6	M83822	Human belge-like protein (BGL) mRNA; partial cds	Other
128131	2.6	AI283162	claudin 3	SS, TM
131564	-	AA491465	ESTs	Other
100279		D42084	Human mRNA for KIAA0094 gene; partial cds	Other
130149		J04031	methylenetetrahydrofolate dehydrogenase (NADP+ dependent); methenyltetrahydrofolate cyclohydrolase; formyltetrahydrofolate synthetase	Other
119888	2.6	W81710	ESTs	SS
126638	2.6	AA649257	ESTs	Other
131672	2.6	X12901	Human mRNA for villin	Other
134405	2.6	J04177	collagen; type XI; alpha 1	Other
101188	2.5	L20320	cyclin-dependent kinase 7 (homolog of Xenopus MO15 cdk-activating kinase)	TM
104534		R22303	ESTs; Weakly similar to putative p150 [H.sapiens]	Other
135179	~ ~	U43747	Friedreich ataxia	SS
124308		H93575	ESTs	SS
125621		AI051602	acetyl-Coenzyme A acetyltransferase 2 (acetoacetyl Coenzyme A thiolase)	TM
101342	a)	L76191	Interleukin-1 receptor-associated kinase 1	Other
129351		AA167268	`ESTs	Other
103774		AA092898	ESTs; Weakly similar to R07G3.8 [C.elegans]	Other
131289		AA485697	ESTs	SS, TM
132094		W01996	ESTs; Highly similar to (define not available 4929683) [H.saplens]	Other
132094	2.5	X74801	chaperonin containing TCP1; subunit 3 (gamma)	Other

FIGURE 11 (CONT) 5 of 10

			5 OJ 10	
PRIMEKEY	Fold Upregulated in Tumors	Exemplar Accession	Complete Titte	ORF Structuaral
105610	2.5	AA279991	ESTs	Other
115719	2.5	AA416997	ESTs	Other
134888	2.5	AA148094	Tat-Interacting protein (30kD)	Other
105564	2.5	AA262943	ESTs	Other
129689	2.5	AA130156	ESTs	Other
103391	2.5	X94453	pyrroline-5-carboxylate synthetase (glutamate gamma-semialdehyde synthetase)	Other
104182	2.5	AA479990	ESTs; Weakly similar to glioma amplified on chromosome 1 protein [H.saplens]	SS, TM
123494	2.5	AA599786	ESTs	Other
122905	2.5	AA470070	ESTs	Other
109175	2.5	AA180496	ESTs	Other
129907	2.5	D80170	ESTs	Other
******	fer e mermene e :		ESTs; Weakly similar to !!!! ALU SUBFAMILY SO WARNING ENTRY !!!!	
115142	2.4	AA258116	[H.sapiens]	Other
125474	2.4	AA151216	Homo sapiens 14-3-3 protein mRNA; complete cds	SS
125745	2.4	AI283493	ribophorin (I	Other
100103	2.4	AF007875	dolichyl-phosphate mannosyltransferase polypeptide 1; catalylic subunit	Other
. 102687	. 2.4	U73379	Human cyclin-selective ubiquilin carrier protein mRNA; complete cds	Other
133170	2.4	U21049	epithelial protein up-regulated in carcinoma	SS, TM
109141	2.4	AA176428	ESTs	Other
132811	2.4	U25435	transcriptional repressor	Other
102862	2.4	X01057	Interleukin 2 receptor, alpha	SS, TM
104300	2.4	D37933	syntaxin 1B	Other
105091	2.4	AA148859	ESTs; Moderately similar to IIII ALU SUBFAMILY J WARNING ENTRY !!!! [H.sapiens]	Other
110674	2.4	H89315	"yw25e09.s1 Morton Fetal Cochlea Homo saplens cDNA clone IMAGE:253288 3', mRNA sequence."	TM
122571	2.4	AA452600	EST	Other
123421	2.4	AA598440	.ESTs; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens]	Other
132181	2.4	AA046939	ESTs; Moderately similar to !!!! ALU SUBFAMILY SX WARNING ENTRY !!!! [H.sapiens]	Other
129445	2.4	AA306121	ESTs	SS ;
113923	2.4	W80763	ESTs; Moderately similar to FK506-binding protein 65kD [M.musculus]	Other
100598	2.4	HG2463-HT2 559	Guanine Nucleotide-Binding Protein G25k	Other
130869	2.4	AA128100	uridine monophosphate synthetase (orotate phosphoribosyl transferase and orotldine-5'-decarboxylase)	Other
132393	2.4	W85888	ESTs; Moderately similar to IIII ALU SUBFAMILY SQ WARNING ENTRY [IIII [H.saplens]	Other
		HG4074-HT4		
100830	2.4	344	-Rad2	Other ;
133765	2.4	D21255	cadherin 11 (OB-cadherin; osteoblast)	SS, TM
100867	2.4	HG4316-HT4 586	Transketolase-Like Protein	Other
119859	2.4	W80702	ESTs	Other
135269	2.4	M77698	YY1 transcription factor	Other
115140	2.4	AA258030	ESTs; Weakly similar to (defline not available 3874821) [C.elegans]	Other
102162	2.4	U18291	CDC16 (cell division cycle 16; S. cerevislae; homolog)	Other

FIGURE 11 (CONT) 6 of 10

PRIMEKEY In Tumors Accession Complete Title	ORF Structuaral Info TM Other Other Other Other SS Other SS Other
100372	TM Other Other Other Other SS Other SS Other Other Other Other Other SS,TM
106981 2.4 AA521157 ESTs 130114 2.4 AA234717 ESTs 116129 2.3 AA439956 ESTs 122235 2.3 AA439475 ESTs 107315 2.3 T62771 Homo sapiens nucleoplasmin-3 (NPM3) mRNA; complete cds draperonin containing TCP1; subumit 2 (beta) 133061 2.3 AB000584 prostate differentiation factor protein phosphatase 1G (formerly 2C); magnesium-dependent; gamma 107531 2.3 Y13936 isoform 107531 2.3 AA456264 ESTs; Highly similar to (defline not available 4176448) [H.sapiens] 101226 2.3 AA456264 ESTs; Highly similar to (defline not available 4809026) [H.sapiens] 101226 2.3 L27706 chaperonin containing TCP1; subumit 8A (zeta 1) 132571 2.3 R84594 ESTs; Highly similar to (defline not available 4809026) [H.sapiens] 100090 2.3 AC002486 saplens] 103658 2.3 Z74615 collagen; type I; alpha 1 104897 2.3 AA456841 ESTs 106818 2.3 AA480890 ESTs 113077 2.3 T40442 ESTs 128773 2.3 M28879 esterase 1) 128773 2.3 M28879 esterase 1) 128773 2.3 M28879 esterase 1) 128773 2.3 AA256286 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 100267 2.3 AA284143 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; complete ds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H.sapiens]	Other Other Other Other SS Other SS Other Other Other Other SS,TM
130114 2.4 AA234717 ESTS 116129 2.3 AA459956 ESTS 122235 2.3 AA436475 ESTS 107315 2.3 T62771 Homo sapiens nucleoplasmin-3 (NPM3) mRNA; complete cds 125905 2.3 T69868 chaperonin containing TCP1; subunit 2 (beta) 133061 2.3 AB000584 prostate differentiation factor protein phosphatase 1G (formerly 2C); magnesium-dependent; gamma soform 107531 2.3 Y13936 soform 125820 2.3 AA730136 teratocarcinoma-derived growth factor 1 131725 2.3 AA456264 ESTs; Highly similar to (defline not available 4176448) [H.sapiens] 101228 2.3 L27706 chaperonin containing TCP1; subunit 8A (zeta 1) 132571 2.3 R84594 ESTs; Highly similar to (defline not available 4809026) [H.sapiens] 100090 2.3 AC002486 SSTs; Highly similar to (defline not available 4809026) [H.sapiens] 103658 2.3 Z74615 collagen; type I; alpha 1 104897 2.3 AA054641 ESTs 106818 2.3 AA480890 ESTS 113077 2.3 T40442 ESTs 128773 2.3 M28879 estarso 1) 129506 2.3 AA258286 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 106267 2.3 AA431873 Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; complete cds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H.sapiens]	Other Other SS Other SS Other Other Other Other SS, TM Other SS, TM
116129 2.3 AA459956 ESTs 122235 2.3 AA436475 ESTs 107315 2.3 T62771 Homo sapiens nucleoplasmin-3 (NPM3) mRNA; complete cds 125905 2.3 T69868 chaperonin containing TCP1; subunit 2 (beta) 133061 2.3 AB000584 prostate differentiation factor 107531 2.3 Y13936 prostate differentiation factor 107531 2.3 Y13936 teratocarcinoma-derived growth factor 1 131725 2.3 AA456264 ESTs; Highly similar to (defline not available 4176448) [H.sapiens] 101226 2.3 L27706 chaperonin containing TCP1; subunit BA (zeta 1) 132571 2.3 R84594 ESTs; Highly similar to (defline not available 4809026) [H.sapiens] 100900 2.3 AC002486 collapse; type I; alpha 1 104897 2.3 AA4564641 ESTs 106818 2.3 AA480890 ESTs 113077 2.3 T40442 ESTs 113077 2.3 T40442 ESTs 128773 2.3 M28879 granzyme B (granzyme 2; cytotoxic T-lymphocyte-associated serine esteraso 1) 128773 2.3 M28879 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 X76040 Lon protease-like protein 106267 2.3 AA431873 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; complete cds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H.sapiens]	Other Other SS Other Other Other Other Other SS, TM Other SS, TM
122235 2.3	Other SS Other SS Other Other Other Other Other SS, TM
107315 2.3 T62771 Homo saplens nucleoplasmin-3 (NPM3) mRNA; complete cds 125905 2.3 T69868 chaperonin containing TCP1; subunit 2 (beta) 133061 2.3 AB000584 prostate differentiation factor protein phosphatase 1G (formerly 2C); magnesium-dependent; gamma isoform 107531 2.3 Y13936 teratocarcinoma-derived growth factor 1 125820 2.3 AA730136 teratocarcinoma-derived growth factor 1 131725; 2.3 AA456264 ESTs; Highly similar to (defline not available 4176448) [H.sapiens] 101228 2.3 L27706 chaperonin containing TCP1; subunit 8A (zeta 1) 132571 2.3 R84594 ESTs; Highly similar to (defline not available 4809026) [H.sapiens] 100900 2.3 R84594 ESTs; Highly similar to (defline not available 4809026) [H.sapiens] 100900 2.3 AC002486 sapiens] 103658 2.3 Z74615 collagen; type I; alpha 1 104897 2.3 AA054641 ESTs 106818 2.3 AA480890 ESTs 113077 2.3 T40442 ESTs 113077 2.3 T40442 ESTs 128773 2.3 M28879 esterase 1) 128506 2.3 AA258286 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 106267 2.3 AA431873 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; complete cds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H.sapiens]	SS Other SS Other Other Other Other Other SS, TM Other SS, TM
125905 2.3 T69868 chaperonin containing TCP1; subunit 2 (beta) 133061 2.3 AB000584 prostate differentiation factor protein phosphatase 1G (formerly 2C); magnesium-dependent; gamma 107531 2.3 Y13936 Isoform 125820 2.3 AA730136 teratocarcinoma-derived growth factor 1 131725; 2.3 AA456264 ESTs; Highly similar to (defline not available 4176448) [H.sapiens] 101228 2.3 L27706 chaperonin containing TCP1; subunit 6A (zeta 1) 132571 2.3 R84594 ESTs; Highly similar to (defline not available 4809026) [H.sapiens] 100090 2.3 R84594 ESTs; Highly similar to (defline not available 4809026) [H.sapiens] 100895 2.3 Z74615 collagen; type I; alpha 1 104897 2.3 AA054641 ESTs 106818 2.3 AA480890 ESTs 113077 2.3 T40442 ESTs 113077 2.3 T40442 ESTs 128773 2.3 M28879 esteraso 1) 129506 2.3 AA258266 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 106267 2.3 AA28143 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; complete cds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H.sapiens]	Other SS Other Other Other Other SS, TM
133061 2.3 AB000584 prostate differentiation factor protein phosphatase 1G (formerly 2C); magnesium-dependent; gamma lsoform 125820 2.3 AA730136 teratocarcinoma-derived growth factor 1 131725; 2.3 AA456264 ESTs; Highly similar to (defline not available 4176448) [H.sapiens] 101228 2.3 L27706 chaperonin containing TCP1; subunit 6A (zeta 1) 132571 2.3 R84594 ESTs; Highly similar to (defline not available 4809026) [H.sapiens] 100090 2.3 AC002486 ESTs; Highly similar to (defline not available 4809026) [H.sapiens] 103658 2.3 Z74615 collagen; type I; alpha 1 104897 2.3 AA054641 ESTs 106818 2.3 AA480890 ESTs 113077 2.3 T40442 ESTs granzyme B (granzyme 2; cytotoxic T-lymphocyte-associated serine esteraso 1) 128773 2.3 M28879 esteraso 1) 129506 2.3 AA258266 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 106267 2.3 AA21873 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; complete cds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H.sapiens]	Other Other Other Other Other Other SS, TM Other SS, TM
protein phosphatase 1G (formerly 2C); magnesium-dependent; gamma 107531 2.3 Y13936 Isoform 125820 2.3 AA730136 teratocarcinoma-derived growth factor 1 131725; 2.3 AA455264 ESTs; Highly similar to (defline not available 4176448) [H.sapiens] 101228 2.3 L27706 chaperonin containing TCP1; subunit 6A (zeta 1) 132571 2.3 R84594 ESTs; Highly similar to (defline not available 4809026) [H.sapiens] "Human BAC clone RG367017 from 7p15-p21, complete sequence [Homo saplens]" 100090 2.3 AC002486 saplens]" 103658 2.3 Z74615 collagen; type I; alpha 1 104897 2.3 AA054641 ESTs 106818 2.3 AA480890 ESTs 113077 2.3 T40442 ESTs 113077 2.3 M28879 esteraso 1) 128773 2.3 M28879 esteraso 1) 129506 2.3 AA258286 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 106267 2.3 AA431873 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; 133493 2.3 AA284143 complete cds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H.sapiens]	Other Other Other Other SS, TM Other SS, TM
107531 2.3	Other Other SS, TM Other SS, TM
125820 2.3	Other Other SS, TM Other SS, TM
131725 2.3	Other SS, TM Other SS, TM
101228 2.3	Other SS, TM Other SS, TM
132571 2.3 R84594 ESTs; Highly similar to (defline not available 4809026) [H.sapiens]	SS, TM Other SS, TM
"Hurman BAC clone RG367O17 from 7p15-p21, complete sequence [Horno sapiens]" 103658 2.3 Z74615 collagen; type I; alpha 1 104897 2.3 AA054641 ESTs 106818 2.3 AA480890 ESTs 113077 2.3 T40442 ESTs granzyme B (granzyme 2; cytotoxic T-lymphocyte-associated serine esteraso 1) 128773 2.3 M28879 esteraso 1) 129506 2.3 AA258266 Horno sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) hornolog 4 134272 2.3 X76040 Lon protease-like protein 106267 2.3 AA431873 Horno sapiens clone 24711 mRNA sequence Horno sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; complete cds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H. sapiens]	Other SS, TM
100890 2.3 AC002486 saplens]* 103658 2.3 Z74615 collagen; type I; alpha 1 104897 2.3 AA054641 ESTs 106818 2.3 AA480890 ESTs 113077 2.3 T40442 ESTs granzyme B (granzyme 2; cytotoxic T-lymphocyte-associated serine esteraso 1) 128773 2.3 M28879 esteraso 1) 129506 2.3 AA258286 Homo saplens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 106267 2.3 AA431873 Homo saplens clone 24711 mRNA sequence Homo saplens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; complete cds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H. saplens]	SS, TM
104897 2.3 AA054641 ESTS 106818 2.3 AA480890 ESTS 113077 2.3 T40442 ESTS granzyme B (granzyme 2; cytotoxic T-lymphocyte-associated serine esterase 1) 128773 2.3 M28879 esterase 1) 129506 2.3 AA258286 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 106267 2.3 AA431873 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; 133493 2.3 AA284143 complete cds 104278 2.3 C02582 ESTs; Highly stmilar to (defline not available 5114045) [H. sapiens]	
106818 2.3 AA480890 ESTs 113077 2.3 T40442 ESTs granzyme B (granzyme 2; cytotoxic T-lymphocyte-associated serine esterase 1) 128773 2.3 M28879 esterase 1) 129506 2.3 AA258286 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 106267 2.3 AA431873 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; complete cds 104278 2.3 C02582 ESTs; Highly stmilar to (defline not available 5114045) [H. sapiens]	Other
113077 2.3 T40442 ESTs granzyme B (granzyme 2; cytotoxic T-lymphocyte-associated serine esterase 1) 128773 2.3 M28879 esterase 1) 129506 2.3 AA258286 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 105267 2.3 AA431873 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 alrophin-1 related protein (DRPLA) mRNA; complete cds 104278 2.3 C02582 ESTs; Highly stmilar to (defline not available 5114045) [H. sapiens]	
granzyme B (granzyme 2; cytotoxic T-lymphocyte-associated serine esterase 1) 129506 2.3 AA258286 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 105267 2.3 AA431873 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; 133493 2.3 AA284143 complete cds 104278 2.3 C02582 ESTs; Highly stmilar to (defline not available 5114045) [H. sapiens]	Other.
128773 2.3 M28879 esterase 1) 129506 2.3 AA258286 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 105267 2.3 AA431873 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 alrophin-1 related protein (DRPLA) mRNA; complete cds 104278 2.3 C02582 ESTs; Highly stmilar to (defline not available 5114045) [H. sapiens]	SS
129506 2.3 AA258286 Homo sapiens mRNA for KIAA0877 protein; partial cds 133746 2.3 U44378 MAD (mothers against decapentaplegic; Drosophila) homolog 4 134272 2.3 X76040 Lon protease-like protein 106267 2.3 AA431873 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; complete cds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H. sapiens]	ss
133746 2.3	TM
134272 2.3 X76040 Lon protease-like protein	Other
105257 2.3 AA431873 Homo sapiens clone 24711 mRNA sequence Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; 133493 2.3 AA284143 complete cds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H. sapiens]	Other
Homo sapiens chromosome 1 atrophin-1 related protein (DRPLA) mRNA; 133493 2.3 AA284143 complete cds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H. sapiens]	Other
133493 2.3 AA2B4143 complete cds 104278 2.3 C02582 ESTs; Highly similar to (defline not available 5114045) [H. sapiens]	
The second secon	Other
THE COLOR AND COMPANY THE PARTY OF THE PARTY	Other
"aa28c03.r1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:814564 127211 2.3 AA480935 5', mRNA sequence."	Other
110721 2.3 H97678 ESTs	Other
114774 2.3 AA150043 ESTs	TM
132968 2.3 N77151 Homo sapiens mRNA for KIAA0799 protein; partial cds	Other
106916 2.3 AA490814 ESTs	Other
113849 2.3 W60439 ESTs; Moderately similar to cbp146 [M.musculus]	Other
131028 2.3 U20240 CCAAT/enhancer binding protein (C/EBP); gamma	Other
Homo sapiens 130 kD Golgl-localized phosphoprotein (GPP130) mRNA;	
130380 2.3 U55853 complete cds	SS
125390 2.3 H95094 KIAA0016 gene product	SS, TM
ESTs; Weakly similar to IIII ALU SUBFAMILY SQ WARNING ENTRY !!!!	
127256 2.3 AA327550 [H.saplens]	Other
132116 2.3 AA234767 ESTs	
117765 2.3 N47797 ESTs	SS, TM
119126 2.3 R45175 ESTs	
129482: 2.2 AA435849 ESTs; Moderately similar to unknown protein [H.sapiens]	SS, TM
120493 2.2 AA255933 ESTs	SS, TM Other
108927 2.2 AA143493 ESTs; Weakly similar to PLECKSTRIN [H.sapiens]	SS, TM Other Other

FIGURE 11 (CONT) 7 of 10

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PRIMEKEY	Fold Upregulated in Tumors	Exemplar Accession	Complete Title	ORF Structuara
····			ESTs; Moderately similar to pregnancy-specific beta-1 glycoprotein 2	
122520	2.2	AA449427	[H.sapiens]	Other
125982	2.2	R98091	RAE1 (RNA export 1; S.pombe) homolog	Other
132325	2.2	N37065	ESTs; Moderately similar to IIII ALU SUBFAMILY SQ WARNING ENTRY IIII [H.sapiens]	Other
100287	2.2	D43950	Human mRNA for KIAA0098 gene; partial cds	Other
114895	2.2	AA236177	Homo sapiens mRNA for KIAA0887 protein; partial cds	Other
105038	2.2	AA130273	ESTs; Weakly similar to (defline not available 4240269) [H.sapiens]	Other
105476	,2.2	AA255473	ESTs	ss
106942	2.2	AA496347	ESTs; Highly similar to gene 7442 protein [H.saplens]	Other
110566	2.2	H59990	ESTs	Other
111068	2.2	N58397	ESTs	Other
127963	2.2	Al299013	gn13h12.x1 NCI_CGAP_Lu5 Homo saplens cDNA done IMAGE:1898183 3' similar to TR:022813 022813 PUTATIVE NAD(P)-DEPENDENT CHOLESTEROL DEHYDROGENASE.;, mRNA sequence.	TM
130985	2.2	AA243700	ESTs	Other
132877	2.2	R40685	Homo saplens mRNA for KIAA0851 protein; complete cds	TM
135242	2.2	M74093	cyclin E1	TM
129468	2.2	J03040	secreted protein; acidic; cysteine-rich (osteonectin)	SS
132616	2.2	AA386264	ESTs	SS, TM
133780	2.2	M14219	decorin	Other
121853	2.2	AA425887	ESTs	Other
114608	2.2	AA079381	ESTs	Other
129017	2,2	H13108	ESTs	SS
132725	2.2	L41887	splicing factor; arginine/serine-rich 7 (35kD)	Other
125606	2.2	R28463	ESTs	SS
105538	2.2	AA258860	ESTs; Highly similar to (defline not available 4583654) [H.sapiens]	Other
133221	2.2	AA235289	ESTs; Highly similar to rap2 gene product (H.saplens)	Other
106389	2.2	AA446949	ESTs	Other
128949	2.2	AA190993	a disintegrin and metalloproteinase domain 12 (meltrin alpha)	Other
120969	2.2	AA398116	ESTs	Other
102495	2.2		and a superior contract the superior of the su	TM
101124	2.2	U51240	Human mRNA for KIAA0085 gene; partial cds	SS
		L10343	protease inhibitor 3; skin-derived (SKALP)	Other
102778	2.2	U83463	syndecan binding protein (syntenin)	
107307	2.2	T52099	creatine kinase; mitochondrial 2 (sarcomeric)	Other
111605	2.2	R11638	ESTs	Other
101959	2.2	\$80343	targinyl-tRNA synthetase	Other
101031	2.2	J05070	matrix metalloproteinase 9 (gelatinase B; 92kD gelatinase; 92kD type IV collagenase)	SS
131701	2.2	AA149008	ESTs; Weakly similar to !!!! ALU SUBFAMILY SQ WARNING ENTRY IIII [H.saplens]	Other
105344	2.2	AA235303	ESTs	TM
103166	2.2	X67951	proliferation-associated gene A (natural killer-enhancing factor A)	Other
131463	2.2	X74142	forkhead (Drosophila)-like 1	Other
131412	2.2	U34044	Human selenium donor protein (selD) mRNA; complete cds	Other
134982	2.2	N46086	ESTs	Other
127236	2.1	AJ341818	budding uninhibited by benzimidazoles 1 (yeast homolog)	Other
10.1			Commence of the Commence of th	

FIGURE 11 (CONT) 8 of 10

			8 0) 10	
PRIMEKEY	Fold Upregulated in Tumors	Exemplar Accession	Complete Title	ORF Structuaral
124596		N70088	ESTs	Other
105650	2.1	AA282347	ESTs; Highly similar to (defline not available 4454694) [H.sapiens]	Other
134917	2.1	X87241	FAT tumor suppressor (Drosophila) homolog	SS
132478	2.1	H20906	Homo sapiens mRNA for KIAA0746 protein; partial cds	SS, TM
134464	2.1	N79354	ESTs; Weakly similar to Rga [D.melanogaster]	Other
111221	2.1	N88869	ESTs	Other
****		HG4557-HT4	A CONTRACTOR OF THE CONTRACTOR	
100892	2.1	962	"Small Nuclear Ribonucleoprotein U1, 1snrp"	Other
117170	2.1	H98153	ESTs	Other
124049	2.1	F10523	primase; polypeptide 2A (58kD)	. Other
131692	2.1	D50914	Human mRNA for KIAA0124 gene; partial cds	TM
103003	2.1	X52003	trefoil factor 1 (breast cancer, estrogen-inducible sequence expressed in)	.: SS .
134085	2.1	U20979	chromatin assembly factor I (150 kDa)	Other
126216	2.1	N23870	ESTs	Other
102721	2.1	U79241	Human clone 23759 mRNA; partial cds	Other
128040	2.1	AA918528	ESTs	Other
101038		J05249	replication protein A2 (32kD)	SS
106060	2.1	AA417287	C-terminal binding protein 2	Other
103622	2.1	Z48042	basic transcription factor 3	Other
127884	2.1	AA768630	ESTs	Other
130911	2.1	W72906	HIRA interacting protein 4 (dnaJ-like)	Other
104260		AF008192	Homo saplens putative GR6 protein (GR6) mRNA; complete cds	Other
120714		AA292689	ESTs	Other
101578		M34423	galactosidase; beta 1	SS
113443		T86158	ESTs	SS
101516		M28249	Accession not listed in Genbank	Other
		14120243	ESTs; Moderately similar to IIII ALU SUBFAMILY SB WARNING ENTRY	
106480	2.1	AA450373	[III] [H.saplens]	Other
111365	2,1	N92659	ESTs	Other
121121	2.1	AA399371	ESTs; Weakly similar to (defline not available 4688925) [H.sapiens]	Other
124768		: R40177	ESTs	Other
102501	2.1	U51586	Human slah binding protein 1 (SlahBP1) mRNA; partial cds	Other
106432		AA448850	ESTs	Other
132812	**************	R48108	ESTs	i Other
125681		AA394176	accessory proteins BAP31/BAP29	TM
130511		L32137	cartilage oligomeric matrix protein	Other
128219		AA978333	ESTs	Other
130962		AA102051	transmembrane 4 superfamily member 6	. SS, TM
101840		M93056	protease Inhibitor 2 (anti-elastase); monocyte/neutrophil	TM
123928		AA621363	EST	Other
	•		ESTs; Moderately similar to IIII ALU SUBFAMILY SC WARNING ENTRY	Other
132073		N57408	IIII [H.sapiens]	Other
101671	•	M61832	S-adenosylhomocysteine hydrolase	Other
107059		AA608545	ESTs	Other
132791		. AA446088	ESTs: Weakly similar to YY1-associated factor 2 [H.saplens]	Other
103131		X65614	S100 calcium-binding protein P	
104791	2.1	AA029046	ESTs	Other

FIGURE 11 (CONT) 9 of 10

	Fold Upregulated	Exemplar		ORF Structuara
PRIMEKEY	in Tumors	Accession	Complete Title	Info
			ESTs; Weakly similar to similar to Yeast hypothetical protein L8167.12 like	Other
135181	2.1	AA821349	[C.elegans]	Other
104334		D82614	ESTs	Other
132103	·	AA025086	ESTs	Other
126743	2.1	AA179253	poly(A)-binding protein-like 1	
100439	2.1	D87453	Human mRNA for KIAA0264 gene; partial cds	Other
114860	2.1	AA235112	ESTs; Moderately similar to similar to murine RNA-binding protein [H.sapiens]	Other
116732	2.1	F13779	ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H.saplens]	Other
131045	2,1	H30863	ESTs; Weakly similar to The KIAA0132 gene product is related to Drosophila melanogaster ring canel protein. [H.sapiens]	Other
133209	2.1	AA114183	ESTs; Moderately similar to glutamate pyruvate transaminase [H.saplens]	Other
101758	9	M77836	Human pyrroline 5-carboxylate reductase mRNA; complete cds	Other
102214	1	U23752	SRY (sex-determining region Y)-box 11	Other
130100		AA425593	ESTs	Other
109799	2.1	F10770	Homo sapiens clone 669 unknown mRNA; complete sequence	Other
130860		U66061	T-cell receptor; beta cluster	SS, TM
130000	. 4.1	000001	ESTs; Highly similar to 73 kDA subunit of cleavage and polyadenylation	
106685	2.1	AA461551	specificity factor [H.sapiens]	TM.
125724	2.1	AA083407	stimulated trans-acting factor (50 kDa)	Other
134258		L28010	heterogeneous nuclear ribonucleoprotein F	Other
129642		R50008	7-dehydrocholesterol reductase	TM
125336		T86823	ESTs	Other
125303		Z39821	ESTs	Other
		HG3521-HT3	Annual months whether the description and the second secon	
100749	2	715	Ras-Related Protein Rap1b	Other
126185	2	Al393989	ESTs	TM
126880	2	AI151406	ESTs	SS, TM
105263	2	AA227926	ESTs	Other
123288	2	AA495836	·EST-	Other
105807	2	AA393803	ESTs	Other
105040	2	AA131047	KIAA0698 gene product	TM
131977	2	F09788	procollagen-proline; 2-oxoglutarale 4-dioxygenase (proline 4-hydroxylase); alpha polypeptide II	SS
117992		N52000	ESTs	SS, TM
134037		X53586	Human mRNA for integrin alpha 6	SS
132744		X54326	glutamyl-prolyl-tRNA synthetase	Other
101581	2	M34996	"Human MHC cell surface glycoprotein (HLA-DQA) mRNA, 3'end"	SS, TM
103576		Z26317	desmoglein 2	SS
131235		AA223284	ESTs; Moderately similar to (define not available 4680395) [H.sapiens]	Other
127910		AA781881	ESTs; Weakly similar to sperm fertilization protein sp56 precursor [M.musculus]	SS
			hypoxanthine phosphoribosyltransferase 1 (Lesch-Nyhan syndrome)	Other
101552		M31642	The state of the s	Other
106318		AA436570	Homo sapiens mRNA for pre-mRNA cleavage factor I subunit	TM
101192		L20859	Human leukemia virus receptor 1 (GLVR1) mRNA; complete cds	SS
104592		R81003	Homo sapiens serine protease mRNA; complete cds	
129095	2	L12350	thrombospondin 2	SS

FIGURE 11 (CONT) 10 of 10

	Fold Upregulated	Exemplar		ORF Structuaral
PRIMEKEY	In Tumors	Accession	Complete Title	 Info
105705	2	AA290767	ESTs	Other
128461	2	M63262	arachidonate 5-lipoxygenase-activating protein	SS, TM

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16074	>10	AA001507	ESTs	other
25047	>10	AA011031	ESTs	other
25082	>10	AA017257	ESTs	other
16490	>10	AA026418	ESTs	other
25179	>10	AA031268	H.sapiens mF	•
25215	16	AA035540	APOLIPOPR	•
25282	>10	AA044825	ESTS	other
16810	>10	AA053636	ESTs	other
16835	>10	AA054438	ESTs	SS,
10747	>10	AA055841	ESTs	other
10748	>10	AA055892	ESTs	other
16993	>10	AA065300	Human mRN/	i
17051	>10	AA070801	ESTs	TM
10840	>10	AA084104	ESTs	other
7296	>10	AA085661	ESTs	other
7325	>10	AA090842	ESTs Weakly	other
17419	>10	AA113349	EST	?
17541	>10	AA127459	ESTs	other
17559	>10	AA128407	ESTs	other
25669	>10	AA128978	ESTs	7
17600	>10	AA130596	ESTs	other .
10992	>10	AA132523	Homo sapien	other
17654	>10	AA133250	ESTs	TM
17734	>10	AA137246	ESTs	other
25801	>10	AA148530	EST - RC AA	TM
25806	>10	AA149007	EST - NO_A	?
11121	>10	AA156359	Human TAR	-
			ESTs	
11160	>10	AA164289		other
25925	>10	AA164494	ESTs Weakly	other
25934	>10	AA165355	Human clone	? .
17987	>10	AA169379	ESTs	other
18008	>10	AA171895	Homo sapiens	TM .
33953	>10	AA173290	Human HOXA	other
33980	>10	AA180223	CAMP-DEPE	
34013	>10	AA190888	EST - RC_AA	other
34014	>10	AA191348	ESTs	other
34015	>10	AA191353	ESTs	TM
34070	>10	AA196549	ESTs	other
18260	>10	AA206801	ESTs	? .
34105	>10	AA207123	ESTs	SS,
34107	>10	AA209469	ESTs	other
18330	>10	AA216722	Human mRNA	TM
18362	>10	AA223912	Ribonuclease	other
18392	>10		ESTs	SS.
34188	>10	AA228030	ESTs	TM
34197	>10		Homo sapiens	
25948	>10	AA234365	Homo sapiens	
25951	>10	AA234556	EST Sapiens	7
11561	>10	AA234556 AA236533		other
			Evi-1	
26059	>10	AA236685	ESTs	other
26100			Human mRNA	
11603			ESTs Highly	other
7785	>10		EST - AA2433	
34372			ESTs	7
26240	>10	AA252282	Human mRNA	. TM

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36506	>10	AA429610	ESTs	othe
38571	>10	AA430726	EST - RC_AA	SS,
38695	>10	AA433910	ESTs	othe
36739	>10	AA435610	EST - RC_AA	?
36753	>10	AA435688	ESTs	othe
36845	>10	AA436198	ESTs	7
13136	>10	AA436560	ESTs	SS,TM
13143	>10	AA436619	ESTs	SS.
36958	>10	AA442060	ESTs	othe
36962	>10	AA442082	ESTs	?
36981	>10	AA442845	EST	7
13237	>10	AA443971	ESTs Weakly	?
13242	>10	AA445994	ESTs	othe
37057	>10	AA446131	ESTs	othe
37068	>10	AA446312	ESTs Weakly	othe
37074	>10	AA446344	ESTs	SS.
37084	>10	AA446486	Homo sapiens	. 7
37135	>10	AA447540	EST	?
37159	>10	AA447714	EST-RC AA	other
37168	>10	AA447772	ESTs	7
37246	>10	AA449311	Homo saplens	other
37310	>10	AA451707	ESTs	other
37453	>10	AA454610	ESTs	?
37456	>10	AA454632	ESTs	TM
27787	>10	AA454660	ESTs	?
37492	>10	AA455248	EST - RC_AA	other
37546	>10	AA456641	ESTS	TM
37601	>10	AA458864	ESTS	other
37611	>10	AA458996	Human signal	SS.TM
37615	>10	AA459101	Human serine	other
37653	>10	AA460017	ESTs	other
37677	>10	AA460530	Homo sapiens	other
37777	>10	AA464860	Homo sapiens	other
8648	>10	AA465016	Homo saplens	?
37816	>10	AA469954	EST	7
37829	>10	AA470084	ESTs	. other
28015	>10	AA477421	ESTs	other
37978	>10	AA479294	EST - RC AA	other
37979	>10	AA479295	ESTs Highly	other
37983	>10	AA479348	H.saplens mR	other
14054	>10	AA485223	ESTs III	TM
38121	>10	AA485724	EST - RC AA	other
28122	>10	AA485928	ESTs Weakly	
38167	>10	AA487207		other
38172		AA487424	1.00	other
38172	>10 >10	AA487424	EST - RC_AA	other
			Homo sapiens	other
38182	>10	AA487501	ESTs	other
38194	>10	AA487969	ESTs	other
20171	>10	AA488432	ESTs	7
38211	>10	AA488687	ESTs	other
38235	>10	AA489030	ESTs	other
38280	>10	AA489791	EST - RC_AA	other
38316	>10	AA490500	Homo saplens	other
38330	>10	AA490882	ESTs	other
38456	>10	AA504343	ESTs	ss,
38460	>10	AA504462	ESTs	other

38553	>10	AA521471	iESTs
38580	>10	AA598545	!ESTs
38590	>10	AA598648	Human mRNA
38601	>10	AA598738	ESTs
28323	>10	AA599639	ESTs
38828	>10	AA609177	
			ESTs
38838	>10	AA609215	EST - RC_AA
38867	>10	AA609318	Human cbl-b r
38871	>10	AA609333	EST
38970	>10	AA609749	ESTs
38984	>10	AA609839	ESTs Modera
39045	>10	AA610077	ESTs
39062	>10	AA620333	EST
39080	>10	AA620552	EST - RC_AA
39110	>10	AA620709	ESTs Weakly
39176	· >10	AA621091	ESTs
39218	>10	AA621330	ESTs
39221	>10	AA621346	Homo sapiens
39232	>10	AA621409	ESTs
21	>10	AB000905	H.sapiens his
8963	>10		TEAFEX-HUMTE
33890	>10		TEAFEX-HUMTE
39302	>10	C14944	ESTs
39329	>10	C20797	EST
28644	>10		
		D12163	ESTs
218	>10	D13540	PROTEIN-TYR
236	>10	D13645	Human mRNA
9127	>10	D30037	PHOSPHATID
459	>10	D38293	Human mRNA
39405	>10	D50975	ESTs
39433	>10	D52037	Human thymid
39436	>10	D52692	Human Ca2+
14708	>10	D59388	EST
39488	>10	D60831	ESTs
39504	>10	D80632	ESTs
765	>10	D86096	Prostaglandin
787	>10	D86969	Human mRNA
789	>10	D86971	Human mRNA
39529	>10	F02202	ESTs
39535	>10	F02450	ESTs Modera
18676	>10	F04022	ESTs
18718	>10	F04915	ESTs
18762	>10	F09458	ESTs
18782	>10	F09739	ESTs
			
29080	>10	F13655	ESTs Modera
19001	>10	H02890	ESTs
19164	>10	H10395	EST
39725	>10	H11323	ESTs
19203	>10	H11593	ESTs
19328	>10	H17808	ESTs .
19387	>10	H20128	ESTs .
39787	>10	H20131	ESTs
	>10	H20165	EST
19389	- 10		
	>10	H26279	EST - RC H2
19389		H26279 H40688	EST - RC_H2

19727	: >10	!H52702	ESTs	~ 1
19787	>10	H56679	ESTs	oth
39995	>10	H62474	EST	SS,T
29331	>10	H68116	ESTs	TM
29344	>10	H68839	EST	નું "
40064	>10	H72283	Human mRN	
40083	>10	H73466	MITOCHON	_;
19949	>10	H78263	ESTs	TM
40204	>10	H88296	EST - RC H	
29523	>10	H88353	ESTs Weak	_
29551	>10	H90134	ESTs	7
29645	>10	H95840	ESTs	7
20057	>10	H98079	ESTs	othe
976	>10		EST - HG203	
1158	>10		EST - HG334	
1210	>10	HG37-HT37		
1346	>10		EST - HG471	
1349	>10		EST - HG474	
1445	>10	J03027	MHC class I	4
1570	>10	K01383	EST - K0138	
1684	>10	L07541	Replication fa	4
1852	>10	L17328	Human FEZ2	
1856	>10	L18920	MELANOMA	1
1863	>10	L19161	TRANSLATIO	1
2070	>10	L37378		i .
1	1	<u> </u>	Homo sapien	
2123	>10 >10	L40396 L41349	Homo sapien	s othe
9723	>10	L41349	Phospholipas	e SS, othe
2188	>10	L47276	ESTs	
2343	>10		EST - L47276	ł
2627	>10	M15353 M29610	Eukaryotic tra	othe TM
2857	>10	M58597	Glycophorin E	
3021	>10	M68941	Fucosyltransf	
3163	>10		Protein tyrosii	
	>10		Cathepsin E	?
3196 20088	>10	M86917	Oxysterol bind	
		N20054	ESTs Weakly	. ?
29795	>10	N20641	ESTs Highly	othe
40427	>10	N21147	ESTs	othe
40444	>10	N22140	ESTs Highly	othe
29893	>10	N23003	ESTs	MT
40498	>10	N26086	Homo sapien	SS,
30190		N33024	ESTs	SS,
	>10	N33264	EST	
30207 20304	>10 >10	N33920	H.saplens mf	othe
20304		N34686	Homo sapiens	?
	>10	N34830	ESTs	other
30265 40594	>10	N35115	ESTs	other
40594	>10	N35388	ESTs	other
40631	>10	N38893	Homo sapiens	other
·	>10	N45124	ESTs	other
40660	>10		NUCLEAR FA	other
30610	>10		EST	?
30617			ESTs	other
30631 30790	>10		EST	?
20564			EST	?
20304	-10	11700443	ESTs	TM

			
40760	>10	N57927	ESTs Weakly
30938	>10	N58561	Cathepsin B
20614	>10	N59230	ESTs
20657	>10	N62889	ESTs
31136	>10	N63512	ESTs Weakly
40827	>10	N64051	Homo sapien
31310	>10	N66831	EST
40876	>10	N67607	Human Rho-a
20791	>10	N68057	Homo sapien:
40905	>10	N68738	ESTs
40911	>10	N69114	H.sapiens mR
40913	>10	N69218	ESTs
31484	>10	N69466	ESTs
31619	>10	N73449	ESTs
41005	>10	N79516	ESTs
31818	>10	N89774	
31872			Homo sapien
	>10	N91109	EST
41040	>10	N91948	ESTs
31944	>10	N93193	ESTs
41065	>10	N93618	ESTs
32034	>10	N98926	ESTs Modera
41107	>10	R01634	ESTs
41163	>10	R08176	ESTs
21238	>10	R08564	Plasminogen-
21240	>10	R08613	ESTs
21412 .	>10	R20670	ESTs
21519	>10	R27975	EST - RC_R2
41381	>10	R42278	H.sapiens mR
32189	>10	R43183	ESTs
32195	>10	R43471	ESTs Weakly
21902	>10	R43822	EST
21946	>10	R44707	ESTs
22072	>10	R49406	ESTs
32240	>10	R50976	Ribonuclease
32258	>10	R55623	ESTs
22258	>10	R56432	ESTs
22282	>10	R59197	ESTs
32277	>10	R61493	Human mRNA
22372	>10	R62831	EST
22400	>10	R64109	ESTs
41593	>10	R64129	ESTS
10233	>10	R71427	
41654	>10	R76437	ESTs Highly
22557			THROMBOXA
	>10	R76722	ESTs
22576	>10	R79111	ESTs
	>10	R79777	EST
41678	>10	R80675	EST
41719	>10	R89260	EST - RC_RB
22793	>10	R96208	ESTs
41752	>10	R97063	ESTs
3375	>10	S50223	HKR-T1
3406	>10	S66896	SQUAMOUS
3522	>10	S80267	Spleen tyrosin
41793	>10	T03887	ESTs
23198	>10	T40530	ESTs Weakly
	>10		

FIGURE 12 (cont.)

SS, other TM other other ? other other other other TM other
TM other

7 other other other other

other other

32635	>10	T61116	ESTs
42177	>10	T79786	ESTs
23623	>10	T84047	ESTs
23662	>10	T86674	ESTs
42242	>10	T89579	Homo saplens
23759	>10	T90313	ESTs
23832	>10	T92018	ESTs
32740	>10	T92950	ESTs
42290	>10	T95105	ESTs
3598	>10	U01157	Glucagon-like
3659	>10	U04313	Protease inhib
	1 >10	U10690	-
3799	4		Human MAGE
3870	>10	U14518	Centromere p
3913	>10	U16261	Human MDA-
4029	>10	U21090	Human DNA p
4157	>10	U28811	Human cystel
4178	>10	U30246	Human burne
15006	>10	U30246	Human bumel
4193	>10	U31116	Human beta-s
4306	>10	U36798	Homo sapiens
4362	>10	U39817	Bloom syndro
4386	>10	U40622	DNA repair pr
4388	>10	U40714	Human tyrosy
4455	>10	U43944	MALATE OXID
4477	>10	U45880	Human IAP-lik
4680	>10	U55766	Human Rev in
4702	>10	U57341	EST - U5734
4713	>10	U57721	Human L-kynu
4787	>10	U61145	Human enhan
4862	>10	U65437	Human homeo
4945	>10	U69108	Homo sapiens
4975	>10	U71088	Human MEKS
4994	>10	U72514	Human C2f m
5002	>10	U72761	Human karyop
5021	>10	U73524	Human putativ
5149	>10	U79716	Human reelin
5214	>10	U83303	H.sapiens mR
5243	>10	U85946	Human brain s
32789	>10	W02779	ESTs Modera
42354	>10	W19346	ESTs Modera
42354	>10	W19346 W40150	
			Homo saplens
33006	>10	W46286	ESTs Weakly
	>10	W45891	ESTs Weakly
33109	>10	W59961	Human mRNA
24197	>10	W67277	ESTs
	>10	W69425	ESTs
	>10	W73883	ESTs
33343	>10	W79834	ESTs Weakly
33377	>10	W81219	ESTs Weakly
42602	>10	W86423	ESTs
33556	>10	W90705	Murine leukern
33616	>10	W93726	Protease Inhib
33666	>10	W95876	ESTs
5510	>10	X05360	Cell division o
5558	>10	X07876	Wingless-type
5603	>10	X14253	Teratocarcino

5619	>10	X14850	HISTONE H2A	-
5623	>10	X14975	T-CELL SURF	?
5692	>10	X17644	G1 to S phase	othe
5789	>10	X54925	Matrix metallo	othe
5799	>10	X55330	Aspartylglucos	SS.
5802	>10	X55544	CYCLIC-AMP	?
5857	>10	X58377	Human mRNa	othe
5960	>10	X63575	ATPase Ca+	TM
5963	>10	X63629	Cadherin 3 (F	SS,T
5986	>10	X64810	(Proprotein con	?
6041	>10	X67155	MITOTIC KIN	othe
6095	>10	X69962	Fragile X mer	othe
6106	>10	X70683	SRY (sex dete	TM
6155	>10	X74331	DNA primase	othe
6167	>10	X74987	Ribonuclease	othe
6188	>10	X76029	NEUROMED	TM
6315	>10	X81889	H.sapiens mR	othe
6382	>10	X85133	H.saplens RB	othe
6384	>10	X85137	Human kinesi	othe
6438	>10	X89398	URACIL-DNA	?
6449	>10	X89986	H.sapiens mR	TM
6478	>10	X91648	H.sapiens mR	SS,TN
6479	>10	X91653	EST - X91653	?
6494	>10	X92689	H.sapiens mR	TM
6713	>10	Y08564	EST - Y08564	?
6790	>10	Y12394	Homo sapiens	othe
24915	>10	YEL003w/	EST - YEL003	?
42773	>10	YEL019c/MN	ASEST - YEL019	?
24545	>10	Z38462	ESTs	othe
33713	>10	Z39427	ESTs	othe
33791	>10	240883	ESTs	othe
42766	>10	Z99394	ESTs Modera	othe
21558	>10	R33112	Human AF-6 m	othe
26718	>10	AA282576	ESTs	?
40113	.9955090946	H78003	ESTs	?
10801	.9879448276	AA069285	ESTs Weakly	.othe
37491	.9513600842	AA455239	ESTs Highly	othe
23900	.9272347693	T95789	ESTs	othe
254	.9198395324	D14657	Human mRNA	othe
6885	.8970927914	Z29331	Ubiquitin-con	othe
29693	.8850766398	H97819	ESTs	SS,
26482	.8765189024	AA262491	ESTs	othe
23123	.8699502035	T25306	EST	?
26525	.8160399123	AA278392	ESTs	othe
13110	.7643356605	AA435840	Homo saplens	other
34863	.7087597628	AA299784	EST	other
39432	.7034550083	D51691	Phosphoribos	?
31312	.6513325388	N66845	ESTs Weakly	?
21112	.6358446349	R01179	IESTs	?
31572	.6254820695	N71294	ESTs	. other
17903	.6221229759	AA160259	EST	?
20747	.6094813734		ESTs	other
4676	9.589223908	U55206	Homo sapiens	TM
34363	.5627081023		Homo saplens	other
39094	9.540768988	AA620636	ESTs	other
3888	.5372000133		Human beta-1	7
	·			

39386 9.5062503.59 D12184 ESTS			For-
4192	39386	19.506250529 D12184	ESTs
4507 9.422674945 L47050			
35606 9.412026255 AA402227 ESTs Moders			
4970 .3649551013 U70862			
19829 .3432151573 H58813 EST 14837 .287858414 T40145 ESTs 17336 .2822148675 AA099585 ESTs 40541 .2532836505 N30160 ESTs 29946 .2487643933 H85434 EST 29943 .1797074262 N24786 ESTs Modera 17997 .1629681314 AA169633 EST 21320 .1243463318 R11673 ESTs 13883 .1178796537 AA476917 ESTs Weakly 30539 .0886887776 N49072 ESTs 26380 .0809559376 AA257012 EST 15888 .0595893607 X95632 Human Abl in 40812 .0012874244 N63419 ESTs 903 .9640387906 D90070 ATL-derived P 22674 .9515777733 R87160 ESTs 40807 .951013226 N62995 TRANSCRIP 15244 .919564497 W00904 ESTs 32296 .8658776567 R67075 Zinc finger pro 18269 .85076626284 H47391 ESTs 41607 8.833925517 R67868 CLEAVAGE S 2548 .8299864699 M25897 Platelet factor 7736 .8279341243 AA232121 Human tyrosy 34490 .7844537272 AA262354 ESTs 39939 .7555031142 H533454 EST -RC_H5 25111 .7232692306 AA020787 ESTs 19662 .850765676 RA599477 ESTs 19652 .8716167279 R36239 EST 39863 .6655982852 H04756 ESTS Highly 1042 .6552112324 HG2510-HT2EST -HG2510 32330 .6361115426 R77776 ESTs 25382 .6239458487 AA059007 ESTS 27074 .5900813076 AA01475 ESTS Weakly 3955 .529890918 U18259 MHC class II 4959 .8555646827 U70322 Human tyrosy 34490 .784453727 AA262354 ESTS 39939 .7555031142 H53454 EST -RC_H5 25111 .7232692306 AA020787 ESTS 25382 .6239458487 AA059007 ESTS 27074 .5900813076 AA01475 ESTS Weakly 3955 .529890918 U18259 MHC class II 4959 .8552646827 U70322 Human transp 2315 .5259185806 M14123 EST - M14123 37253 .4896914632 AA449357 ESTS 39624 .471316977 F10836 ESTS 39934 .7555033144 R07499 ESTS 32479 .4093689545 T16262 WEE1-LIKE P 41251 .3587565415 R04766 ESTS Weakly 3925 .529809818 U18259 MHC class II 4959 .855596435 M54995 Connective Its 41154 .441339014 R07499 ESTS 32479 .4093689545 T16282 WEE1-LIKE P 41251 .3587565416 R07499 ESTS 3213 .4569920887 T40891 ESTS 321479 .4093689545 T16262 WEE1-LIKE P 41251 .3587565416 R06965 ESTS 3215 .5259185806 H64966 Human CENF	h		
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17336		.3432151573H58813	
40541 .2532836505 N30160 ESTs			
29496	17336		5 ESTs
29943 1.797074262 N24786 ESTs Modera 17997 1.629681314 AA169633 EST 21320 1.243463318 R11673 ESTs 13883 1.178796537 AA476917 ESTs Weakly 30539 0.686687776 N49072 ESTs 32778 0.677919545 W02063 EST 26380 0.609559378 AA257012 EST 15888 0.595893607 X95632 Human Abl in 40812 0.012874244 N63419 ESTs 903 9640387906 D90070 ATL-derived P 22674 9515777733 R87160 ESTs 40807 951013228 N62995 TRANSCRIP 15244 919564497 W00904 ESTs 32296 .8656776567 R67075 Zinc finge pro 18269 .85576656766 AA209467 ESTs 19662 .8507626284 H47391 ESTs 41607 8.833925517 R67868 CLEAVAGE S 2548 .8299864699 M25897 Platelet factor 7736 .8279341243 AA232121 Human tyrosy 34490 .7844537272 AA262354 ESTs 38658 .7669313482 AA599477 ESTs 7528 8.765157554 AA149543 ESTs 39939 .7555031142 H53454 EST - RC_H5 25111 .7232692303 AA020787 ESTs 21655 8.716167279 R36239 EST 39663 8.665982852 H04756 ESTs Highly 1042 8.652112324 HG2510-HT2 EST - HG2510 32330 .6361115426 R77776 ESTs 25382 .6239458467 AA059007 ESTs 27074 .5900813076 AA401475 ESTs Weakly 3955 .529890918 U18259 MHC class II 4959 8.52646827 U70322 Human transp 2315 .5259185806 M4123 EST - M14123 37253 .4896914632 AA449357 ESTs 39624 8.471316977 F10836 ESTs 32479 .4093689545 T16262 WEE1-LIKE P 41251 .358756541 R28279 Human clone 19081 .3583603163 H06701 ESTs Weakly 21098 .3105927555 R00545 ESTs 37154 .299482234 AA447666 Human CENF	40541	.2532836505N30160	ESTs
17997	29496	.2487643833H85434	EST
21320	29943	.1797074262N24786	ESTs Modera
13883	17997	.1629681314AA16963	3 EST
30539	21320	.1243463318R11673	ESTs
32778	13883	.1178796537 AA47691	7 ESTs Weakly
32778	30539	.0886887776N49072	ESTs
26380			
15888			1
40812 .001287424 N63419 ESTs 903 .9640387906 D90070 ATL-derived P 22674 .9515777733 R87160 ESTs 40807 .951013226 N62995 TRANSCRIP 15244 .9195644974 W00904 ESTs 32296 .8658776567 R67075 Zinc finger pro 18269 .8575656766 AA209467 ESTs 19862 .8507626284 H47391 ESTs 41607 8.833925517 R67868 CLEAVAGE \$ 2548 .8299864699 M25897 Platelet factor 7736 .8279341243 AA232121 Human tyrosy 34490 .7844537272 AA262354 ESTs 38658 .7669313482 AA599477 ESTs 7528 8.765157554 AA149543 ESTs 39939 .7555031142 H53454 EST - RC_H5 25111 .7232692306 AA020787 ESTs 39663 8.665982852 H04756 ESTs Highly 1042 8.652112324 HG2510-HT2 EST - HG2510 32330 .6361115426 R77776 ESTs 27074 .5900813076 AA401475 ESTs Weakly 3955 .5298909183 U18259 MHC class II 4959 8.52646827 U70322 Human transp 2315 .5259185808 M14123 EST - M14123 37253 .4896914632 AA449357 ESTs 39624 8.471316977 F10836 ESTs 23213 .456992088 T40891 ESTs 232479 .409368954 T16262 WEE1-LIKE P 41251 .358756541 R28279 Human cone			
903		1	
22674			
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7528 8.765157554 AA149543 ESTs 39939 .7555031142 H53454 EST - RC_H5 25111 .7232692309 AA020787 ESTs 21655 8.716167279 R38239 EST 39663 8.665982852 H04756 ESTs Highly 1042 8.6552112324 HG2510-H72 EST - HG2510 32330 .636111542 R77776 ESTs 25382 .6239456487 AA059007 ESTs 27074 .5900813076 AA401475 ESTs Weakly 3955 .529890918 U18259 MHC class II 4959 8.52646827 U70322 Human transp 2315 .5259185808 M14123 EST - M14123 37253 .4896914632 AA449357 ESTs 39624 8.471316977 F10836 ESTs 23213 .4569920887 T40891 ESTs 23213 .4569920887 T40891 ESTs 32479 .4093689549 T16282 WEE1-LIKE R 41251 .3587565416 R28279 Human clone 19081 .3563603163 H06701 ESTS Weakly 21098 .3105927555 R00545 ESTS 14723 .3061679053 D59894 ESTS 14723 .3061679053 D59894 ESTS	34490		
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25111723269230\$ AA020787 ESTS 21655 8.716167279 R38239 EST 39863 8.665982852 H04756 ESTS Highly 1042 8.6552112324 HG2510-HT2 EST - HG2510 32330 .6361115426 R77776 ESTS 25382 .6239456487 AA059007 ESTS 27074 .5900813076 AA401475 ESTS Weakly 3955 .5298909183 U18259 MHC class II 4959 8.52646827 U70322 Human trans; 2315 .5259185806 M14123 EST - M14123 37253 .4896914632 AA449357 ESTS 39624 8.471316877 F10836 EST6 23213 .456992086 T40691 ESTS 2798 8.455596435 M54995 Connective IIS 41154 .441339014 R07499 ESTS 32479 .409366954\$ T16262 WEE1-LIKE R 41251 .3587565416 R28279 Human clone 19081 .3583603183 H06701 ESTS Weakly 21098 .3105927555 R00545 ESTS 37154 .299482234 AA447666 Human CENR	7528	8.765157554 AA14954	3 ESTs
21655 8.716167279 R36239 EST 39663 8.665982852 H04756 ESTS Highly 1042 8.652112324 HG2510-HT2ÆST - HG2510 32330 .6361115426 R77776 ESTS 25382 .6239456487 AA059007 ESTS 27074 .5900813076 AA401475 ESTS Weakly 3955 .5298909183 U18259 MHC class II 4959 8.52646827 U70322 Human transp 2315 .5259185806 M14123 EST - M14128 37253 .4896914632 AA449357 ESTS 39624 8.471316877 F10836 ESTS 23213 .4569920887740891 ESTS 2798 8.455596435 M54995 Connective IIS 41154 .441339014 (R07499 ESTS 32479 .4093689549 T16282 WEE1-LIKE R 41251 .3587565416 R26279 Human clone 19081 .358360318 106701 ESTS Weakly 21098 .3105927559 R00545 ESTS 14723 .3061679053 D59894 ESTS 37154 .299482234 (AA447666 Human CENF	39939	.7555031142H53454	EST - RC_HS
39863 8.665982852 H04756 ESTs Highly 1042 8.652112324 HG2510-HT2EST - HG2510 32330 .6361115426 R77776 ESTs 25382 .6239458487 AA059007 ESTs 27074 .5900813076 AA401475 ESTs Weakly 3955 .5299909183 U18259 MHC class II 4959 8.52646827 U70322 Human transp 2315 .5259185806 M14123 EST - M14128 37253 .4896914632 AA449357 ESTs 39624 8.471316877 F10836 ESTs 23213 .4569920887 T40891 ESTs 2799 8.455596435 M54995 Connective IIs 41154 .441339014 (R07499 ESTs 32479 .4093689549 T16282 WEE1-LIKE R 41251 .3587565416 R26279 Human clone 19081 .3583603183 H06701 ESTs Weakly 21098 .3105927559 R00545 ESTs 14723 .3061679053 D59894 ESTS 37154 .299482234 (AA447666 Human CENR	25111	.7232692309 AA02078	7 ESTs
1042 8.652112324 HG2510-HT2 EST - HG2510 32330 .6361115426 R77776 ESTs 25382 .6239456467 AA059007 ESTs 27074 .5900813076 AA401475 ESTs Weakly 3955 .5299909183 U18259 MHC class II 4959 8.52646827 U70322 Human transp 2315 .5259185808 M14123 EST - M14123 37253 .4896914632 AA449357 ESTs 39624 8.471316877 F10836 ESTs 23213 .4569920887 T40891 ESTs 2799 8.455596435 M54995 Connective IIS 41154 .441339014 (R07499 ESTs 32479 .4093689549 T16282 WEE1-LIKE R 41251 .3587565416 R26279 Human clone 19081 .3583603183 H06701 ESTs Weakly 21098 .3105927559 R00545 ESTs 14723 .3061679053 D59894 ESTS 37154 .299482234 (AA447666 Human CENF	21655	8.716167279 R38239	EST
1042 8.652112324 HG2510-HT2 EST - HG2510 32330 .6361115426 R77776 ESTs 25382 .6239456467 AA059007 ESTs 27074 .5900813076 AA401475 ESTs Weakly 3955 .5299909183 U18259 MHC class II 4959 8.52646827 U70322 Human transp 2315 .5259185808 M14123 EST - M14123 37253 .4896914632 AA449357 ESTs 39624 8.471316877 F10836 ESTs 23213 .4569920887 T40891 ESTs 2799 8.455596435 M54995 Connective IIS 41154 .441339014 (R07499 ESTs 32479 .4093689549 T16282 WEE1-LIKE R 41251 .3587565416 R26279 Human clone 19081 .3583603183 H06701 ESTs Weakly 21098 .3105927559 R00545 ESTs 14723 .3061679053 D59894 ESTS 37154 .299482234 (AA447666 Human CENF	39863	8.665982852 H04756	ESTs Highly
25382 .6239456487 AA059007 ESTs 27074 .5900813076 AA401475 ESTs Weakly 3955 .5298909183 U18259 MHC class II 4959	1042	8.652112324 HG2510-I	
27074 .5900813076 AA401475 ESTs Weakly, 3955 .5298909183 U18259 MHC class II 1 4959	32330	.6361115426R77776	ESTs
27074 .5900813076 AA401475 ESTs Weakly, 3955 .5298909183 U18259 MHC class II 1 4959			
3955 .529890918. U18259 MHC class II 4959 8.52646827 U70322 Human transp 2315 .5259185808 M14123 EST - M14128 37253 .4896914632 AA449357 ESTs 39624 8.471316877 F10836 ESTs 23213 .4569920867740891 ESTs 2798 6.455596435 M54995 Connective II3 41154 .441339014 (R07499 ESTs 32479 .4093689549716282 WEE1-LIKE R 41251 .3587565415 R28279 Human clone 19081 .3583603183 H06701 ESTs Weakly, 21098 .3105927559 R00545 ESTs 14723 .3061679053 D59894 ESTs 37154 .299482234 (AA447666 Human CENF			
4959 8.52646827 U70322 Human transp 2315 .5259185806 M14123 EST - M14123 37253 .4896914632 AA449357 ESTs 39624 8.471316877 F10836 ESTs 23213 .4569920887T40891 ESTs 2709 6.455596435 M54995 Connective Its 41154 .441339014 R07499 ESTs 32479 .4093689549 T16282 WEE1-LIKE R 41251 .3587565418 R28279 Human clone 19081 .358360318 H06701 ESTs 21098 .3105927558 R00545 ESTs 14723 .3061679053 D59894 ESTs 37154 .299482234 AA447666 Human CENF			
2315 .5259185808 M14123 EST - M14123 37253 .4896914632 AA449357 ESTs 39624 8.471316977 F10836 ESTs 23213 .4569920887 T40891 ESTs 2798 8.455596435 M54995 Connective tis 41154 .441339014 (R07499 ESTs 32479 .4093689549 T18282 WEE1-LIKE R 41251 .3587565415 R28279 Human ctone 19081 .3583603183 H06701 ESTs Weakly 21098 .3105927555 R00545 ESTs 14723 .3061679053 D59894 ESTS 37154 .299482234 (AA447666 Human CENR			
37253			
39624 8.471316877 F10836 ESTs 23213 .4569920887 T40891 ESTs 2798 8.455596435 M54995 Connective tis 41154 .4413390141 R07499 ESTs 32479 .4093689549 T16282 WEE1-LIKE F 41251 .358756541 R28279 Human clone 19081 .3583603183 H06701 ESTs Weakly 21098 .3105927559 R00545 ESTs 14723 .3081679053 D59894 ESTs 37154 .299482234 AA447666 Human CENF			
23213			
2798 6.455596435 M54995 Connective its 41154 .4413390141 R07499 ESTs 32479 .4093689549 T18282 WEE1-LIKE R 41251 .358756541 R28279 Human done 19081 .3583603183 H06701 ESTs Weakly 21098 .3105927558 R00545 ESTs 14723 .3061679053 D59894 ESTs 37154 .299482234 AA447666 Human CENP			
41154 441339014 R07499 ESTs 32479 4093689549 T18282 WEE1-LIKE R 41251 3587565416 R28279 Human clone 19081 3583603183 H06701 ESTs Weakly 21098 3105927559 R00545 ESTs 14723 3061679053 D59894 ESTs 37154 299482234 AA447666 Human CENR	·		
32479 .409368954\$T16282 WEE1-LIKE R 41251 .358756541\$R28279 Human clone 19081 .3563603163.H06701 ESTs Weakly 21098 .310592755\$R00545 ESTs 14723 .3061679053.D59894 ESTs 37154 .299482234 AA447666 Human CENR			
41251 .358756541ER28279 Human clone 19081 .3583603183H06701 ESTs Weakly 21098 .310592755ER00545 ESTs 14723 .3061679053D59894 ESTs 37154 .299482234 AA447666 Human CENF			
19081 3583603183 H06701 ESTs Weakly 21098 3105927555 R00545 ESTs 14723 306167905 D59894 ESTs 37154 299482234 AA447666 Human CENP			
21098 .3105927558R00545 ESTs 14723 .3061679053D59894 ESTs 37154 .299482234 AA447666 Human CENP			
14723 .3061679053 D59894 ESTs 37154 .299482234 AA447666 Human CENP			
37154 .299482234 AA447666 Human CENF			
8068 .283558636 AA313387 ESTs Highly			
	8068	.2835586361AA313387	/ ESTs Highly

FIGURE 12 (cont.)

TM other TM TM other ? TM other other ? TM other
other other TM other TM other other

other

other other

other ? other ? other other SS, other

тм

other other other other

7485	8.281679348	ΔΔ129547	ESTs	other
16501	.251796983		ESTs	other
34527	.241916375		ESTs	other
6700	.194867566		H.sapiens mR	other
2852	.192881653	1	Human 75-kD	other
11188	.186249246		ESTs	TM
42293	8.183311064		ESTs Weakly	TM
5443	.176331754		Interferon (gan	
40937	.153481059		ESTS	TM
23371	.149949606		EST - RC T59	
26272	.1339974519		ESTs Weakly	other
17306	.1332403762		ESTs	other
18497	.1192326373		ESTs	other
235	.094436390		Human mRNA	other
24525	.0860187097		ESTs	TM
7826	.0750029554		EST - AA2488	TM
32142	.0739258775		Homo sapiens	other
39067	.0557768803		ESTs	other
6235	.0448957236		Casein alpha	TM
29517	.0017588725		ESTs	other
28570	.9852455973	<u> </u>	Homo sapiens	other
39344	.9162087762		ESTs Modera	other
18951	.9002189759		ESTS	other
18953	.8709160227		ESTs	other
18376	.8564099916		ESTs	other
19830	7.847878447		ESTs	other
36023	7.840835828		ESTS	other
13347	.8344414518		ESTs	other
36614	.8284591351		ESTs	other
2192	.8254072032		Homo Sapien	?
33016	.8006574068		H.saplens mR	other
17215	.7941954038		ESTs	other
34894	.7659738105		EST	?
40614	7.695001222		ESTs	other
36295	.6834749899		ESTs	other
19564	.6744302788		ESTs	· TM
16914	.6686405336		ESTs	SS.
35967	.6378079107		Human splicin	other
21672	.6364823402		ESTs	other
19918	.6303275831		ESTs	7
10511	.6297744492		ESTs Highly	other
17721	.6057911016		ESTs	?
42302	.6031859697		EST	SS.
26134	.6000619383		ESTs	other
18766	.5621799008		ESTs	other
34492	7.501590494		ATL-derived P	other
270	.4512152125		EST - D14822	other
35975	.4177746986		ESTs	other
29842	.4095809671		ESTs	7
35389	.3913043319		ESTs	other
19979	.3868157166		ESTs	other
5793	.3885864025		CDC28 protei	other
19978	7.380969715		EST - RC HE	other
1280			EST - HG4126	7
31571	.3676263454		ESTs	other
23765	.3541191734		ESTs Weakly	7

35123	339793345	AA380927	IEST	?
38252	.3341119467	AA489247	ESTs	other
38216	.3282021037	AA488861	ESTs	other
29418	.2489407005	H77915	EST - RC H7	7
4834	.1980951054	U63541	Human mRNA	. other
42504	.1913036522	W69803	ESTs	other
6111	7.158000198		H.saplens mR	TM
41773	7.154479618		ESTs Weakly	other
9951	.1363626365		ESTs	other
28109	.0941968224		ESTs	other
988		<u> </u>	EST - HG2160	7
29848	.0610668511		ESTs	other
30628	.0607950168		ESTs	other
22567	.0225726353		ESTs	TM
9347	7.006323071		ESTs	TM
11696	.0026773299		ESTs	other
40584	.0010096333		EST	?
193	.9767029188		PROBABLE G	TM
18305	.9740536051		Collagen type	other
6078	.9699682397		FARNESYL-D	other
26741	6.902658703		ESTs	other
35069	.8992865685		EST	7
23504	.8977135983		ESTs	other
299	.8824513029		Homo sapiens	other
40583	.8689903023		ESTs	other
31428	.8623762224		ESTs	other
6169	.8606959727		SET PROTEIN	
39524	.8587355171		MALATE OXID	
34578			ESTs UNID	other
38678	.8430889439 6.837527995		Small inducible	other
23936				other
9326	.8251471804		ESTs	other
19188	.8181321394 .8067351968		Msh (Drosoph	TM
18185	.7882148811		ESTs Highly	other
27028			Homo sapiens	
41289	6.757529124 .7519531681		ESTs Weakly	other
34511			EST DO A	
	.7364448798		EST - RC_AA	other ?
1566	.7056207716		EST - J05614	•
25675	.6692299748		ESTs Highly	other
5814 13861	.6584342828		CYTOCHROM	SS,
	.6236291607		ESTs	other
29794 39333	.6026313352		ESTs	other
	.5902382643		Cyclin B1	other
3770	.5835303599		Nuclear factor	other
31831	.5829933764		ESTs	?
33063	.5808125026		Homo sapiens	other
20326	.5640084836		ESTs Weakly	7
34384	.5535703492		ESTs	other
25599	.5490481991		Human (clone	other
39749	.5369363254		ESTs	other
42596	.5200567072		ESTs	?
39606	.5119482185		ESTs Weakly	7
14617	.5105504748		ESTs	other
27831	6.45670814		ESTs	?
34896	.4496517783		EST	7
27360	.4434305008	AA425356	ESTs	other

FIGURE 12 (cont.)

20126	.4326610424N22015	ESTs	тм
6663	.4324809977Y00291	RETINOIC AC	TM
30692	.4196636207N51563	ESTs	other
36472	.4189542265 AA428633	EST	7
9578	.3961788753H87652	Homo saplens	other
39670	.3818496159H05626	ESTs	other
22697	.3652792447 R89218	ESTs	other
37308	.3647804993 AA451694	EST	TM
16101	.3517262802 AA002147	EST	7
20629	.3486854401N59798	ESTs	other
36100	.3364146287 AA417740	ESTs	?
15488	.3252590241W28097	Homo sapiens	other
36667	.3131273544AA432136	ESTs	other
30766	.3115037924N52627	EST - RC_NS	7.
32882	.2745311453W37683	ESTS	TM
18072	.2675797205 AA180448	EST	7
18231	.2652604863 AA199747	Human mRNA	other
38282	.2514165678 AA489814	EST	?
28125	6.250317021 AA486073	ESTs	other
37464	.2484456382 AA454747	ESTs	7
36618	.1946328223 AA431478	ESTs	other
5082	.1931116815U78524	Human Gu bin	other
1441	.1777287039 J02963	Integrin alpha	other
42105	6.14875944 T67710	ESTs	?
6061	.1394863141X68314	Glutathione pe S	ss,
32570	.1156028796 T30222	ESTs Weakly	TM
32504	.1019612076T17063	EST	?
23335	.0977927504 T56804	EST	?
10867	.0970991075 AA088458	ESTs Weakly	other
30883	.0911993489N56923	EST	?
14528	.0859008453 AA620295	ESTs	TM
29454	.0685955036H81308	EST	?
6798	.0539173278 Y13153	Homo sapiens	TM
21248	.0525426545 R08871	ESTs	?
21940	.0499964138R44538	ESTs	?
29066	.0455247653F10927	Homo sapiens	other
18774	.0446826953F09609	ESTs	?
36722	.0172343991 AA435512	120.0	SS,
18062	.0034342969 AA179845	1.00.0	other
22989	.9992817406 T16305	1	other
41745	.9905623898 R95895	ESTs	7
8787	.9894877658 AA504307	77. 07. 11. 11. 1	other
20550	5.984861795 N55013	1	other
26470	.9417764101AA262179		other
16574	.9356497569 AA031926		other
693	.9169537385 D80007		other
4093	5.914830973 U25182	Trainer, and	TM
1192	.9086264407 HG3546-HT		?
22956	1.8954735623T10248		other
36723	5.891606409 AA435524	EST	?
2114	1.8844986595L40384		other
26872	5.868238789 AA291137	100.0	other
6602	.8663883018 X98266	4	other
42701	.8594493433Z38612	1	other
28573	5.84591116 C21118		other other
18290	1.8189427595 AA211901	ESTs	Ulitef

FIGURE 12 (cont.)

732	.8043917941	D83781	Human mRNA	other
5330	.8014145611	U91327	EST - U91327	?
33503	.7990715189	W88720	EST	?
2553	.7797505864	M26167	Human platele	?
34705	.7658806254	AA286907	ESTs Weakly	other
42665	.7594091043	W93659	ESTs	other
38180	.7539310793		EST - RC_AA	other
4244	.7476738809	U33286	Human chrom	other
32822	.7418957453	W16834	ESTs	TM
3977	.7245885557	U18991	Retinal pigme	?
24673	.7202366155	Z39301	ESTs	TM
6928	.7120261126	Z46629	SRY (sex-dete	other
38726	.7030796258	AA608733	ESTs	?
39290	.6892372058		Human mRNA	other
11405	.6818873796		ESTs	other
22538	.6792006591	R73567	Homo sapiens	TM
40747	.6605393208		Homo saplens	TM
31596	.6554024604		ESTs	other
6329	.6415652518		EST - X82279	?
31578	.6273323661		ESTs	other
33207	.6271818482		H.saplens mR	other
2545	.6105860146		Cyclin B1	other
22580	.5988402647		ESTs	other
33592	.5935314518		ESTs	other
28843	.5734698755		ESTs	other
6160	.5689050619		CDC21 HOMO	other
37987	5.561345667		ESTs	other
42515	.5217868611		Homo sapiens	other
4732	.5130688527		Human huntin	other
3299	.5099850678		Hydroxymethy	?
28320	5.473406981		ESTs	?
746	5.471260899		Human mRNA	TM.
39373	.4635804954		ESTs	other
3117	.4398413537		Peroxisomal m	other
21257	.4343612441		ESTs Modera	other
31487	.4318648859		ESTS	other
28954	.4137130511		ESTs	other
38928	5.389782721		ESTs	other
29903	.3722320622		EST	7.
30925	.3437432315		ESTs Weakly	7
19091	.3344615669		ESTs	TM.
28209	.3138951918		ESTs	other
9470	.3118897984		EST - H46617	other
9435	.3070056656		EST - H30201	7
28552	.2954432572		ESTs	other
27411	.2934432572		ESTs	other
30615	.2924125264		ESTs	other
28313	.2657977167		ESTS	TM
39321	.2649035384		ESTs	1M
29934	.2531047395		ESTs	other
1094			EST - HG284B	7
39578	.2481126384		ESTs	TM
11232	.2461120304		ESTs Weakly	other
2466	.2426349328		Human small	other
26843	.2387758661		ESTs	Olliei Olliei
40331	.2353385567		ESTs Weakly.	r other
40331	.2333303301	1.131304	LOTO TEARIY.	,

8035	5.205798365		EST - AA305	1
29793	.195542572		ESTs Weakly	othe
34109	.148159010	AA210722	EST	?
26408	.143257725	AA258177	ESTs Weakly	othe
19263	.1427029807	H15054	ESTs	TM
24596	.1416089352	Z38810	ESTs	othe
28589	.1365059753	C21245	H.saplens mF	othe
5684	.1121931412		Pregnancy-sp	othe
30710	.1079347344		EST	?
35765	.0973514948		EST	7
26360	.086312786		ESTs	.?
	.0849612092		Proliferating of	?
2351	.0836877534			
30262			Homo sapiens	
41792	.0737512465		ESTs	?
36710	.0703839864		ESTs	othe
39090	.0546885407		ESTs	TM
42185	.0539926381		ESTs	?
18745	.0460321557		ESTs	other
35746	.0396841996		ESTs	other
35356	.0354809581		EST	?
36769	.0312706878		EST	7
36900	.0279911548	AA436866	H.saplens mf	other
27595	.0244757301	AA443328	ESTs	TM
16290	.0056611904	AA016145	ESTs	?
27117	.0016146599	AA405098	ESTs Weakly	other
4304	.9951954397	U36764	Eukaryotic tra	other
33458	.9907402071	W86835	Homo sapien	other
26693	.9800090679	AA282120	EST	?
12669	.9758138651	AA417030	Homo sapien	other
29701	.9708526387	H97970	EST	7
20480	.9557253636	N52168	ESTs	TM
8720	.9439110602	AA481218	EST - AA481	e other
34828	.9431269475		Homo sapiens	SS,TM
14985	4.941621032	U15128	Human beta-	?
16115	.9377553522		ESTs	7
42506	.9348587118		EST	other
34761	.9316837445		ESTs	other
11870	.9281056201		ESTs	TM
23211	.9258391854		ESTs	other
40611	.9160502275		Homo sapiens	other
42611	.9128605354		Homo sapiens	other
39652	.9045174605		ESTs	other
17581	4.889674751		EST	?
37239	.8704375389		ESTs	,
18712	.8703618781		ESTs	other
30709	.8611171953		ESTs Weakly	other
34179	.8503613948		ESTs Highly	other
21433	4.825670988		EST	?
39731	.8186142741		ESTs	r other
	.8116614274		ESTS	other
31295	4.804163055			otner
24647	 		EST	
31292	.8008871817		ESTs	other
1285			EST - HG415	7 ?
1106			EST - HG298	
18212	.7912262565		ESTs	other
34367	4.782207045	JAA251758	Homo sapiens	other

477 1 700 000000000000000000000000000000		
34802	.7797760205AA291468	ESTS TA
34762	.7775301546AA287834	ESTs of
11595	.7696612848 AA242819	ESTs of
8295	.7639839111AA405082	ESTs
17622	4.758635576 AA131584	ESTs Weakly of
35781	.7572463523 AA406335	ESTs oth
34754	.7483874972AA287642	Human mRNA oth
23237	.7444854356T47291	EST
37667	.7280445357AA460318	ESTs Highly oth
11568	.7257189975 AA236786	ESTs oth
38622	.7190695733AA598967	ESTs
5137	7057359474U79296	Dihydrolipoard oth
25038	.7002244728 AA010065	CDC28 protei oth
19288	.7000147312H16567	ESTs oth
32503	.6979488292T17045	Collagen type oth
3278	.6953739298 M94055	SODIUM CHA TM
9696	.6942061018L38961	Integral transm TN
35400	.6901390898 AA399591	Homo saplens oth
35246	.6862691303AA398367	EST Weakly \$
36387	.682249927 AA426270	ESTs oth
21509	.6730072542 R27314	ESTs oth
31381	.6729672124N67889	ESTs oth
26723	.6727894925 AA282781	ESTs Highly oth
36326	.6703621086AA425151	Human GAP S oth
17409	.6688418667 AA113136	EST-RC AA oth
4908	.6552339935U67156	Human mitoge other
30594	.6496238328 N49967	ESTs other
38286	4.64639735 AA489847	ESTs Weakly 7
13073	.6426509459 AA433950	ESTS other
40435	.6240181066N21614	Homo sapiens other
14474	.6228694379 AA609427	ESTs Modera othe
38213	4.615309907 AA488847	ESTs Weakly ?
5312	4.606644198 U90716	Human cell su SS.TI
24225	.6041550359W70326	ESTs 7
35588	.5868982366 AA401750	EST ?
29739	.5863199051H99626	EST ?
7203	.5792992577AA053096	EST - AA0530 othe
2157	.5772055869L41939	Homo sapiens SS.TI
32086	.5661024279R11510	ESTs ?
8085	.5648114738AA314779	ESTs Weakly, SS,
224	.5622018989D13633	
34006	.5609980241AA188761	
33656	.5557384389W95477	DNA polymera othe
34065	.5537335124 AA195517	ESTs othe
6028	1.5357922097X66503	144
		Adenylosuccin othe
4166	.503293067 U29463	Cytochrome B ?
40262	.5024727522 H93562	ESTs TM
22687	.5018672549 R88209	ESTs TM
41069	.4977510482 N93969	H.sapiens mR SS.
8264	.4793100575 AA401334	ESTs othe
	4.472017297 AA443187	ESTs ' othe
35882	.4717597552 AA412047	!ESTs ?
	4.465519191 AA262080	Human burnet TM
15921	.4548516436 Y12065	Homo sapiens ?
11279	.438003867 AA195399	ESTs othe
39222	.4367650786AA621348	ESTs Highly othe

			,
34428	.4364736766		ESTs
8771	4.432067373		ESTs
22193	.4189610024		Homo saplen:
7898	.4066170874		ESTs
19902	.3886145805	H66736	ESTs
9276	3868095209	D82374	ESTs
10716	.3794529068	AA053319	ESTs
13193	.3751913512	AA442763	ESTs Highly
5690	.3723059417		NUCLEOSIDE
35102	4.37147138		EST - RC AA
17983	.3612985467		ESTs
24962		AFFX-HUMTI	
31680	.3416539669		ESTs
27168	4.330306894		ESTs
	1		
28731	.3231848659		EST
28348	.3212284906		ESTs
16335	.3019961487		ESTs Weakly
33036	.2915644973		ESTs Weakly
30180	.2897721925		ESTs
35591	.2895541242		ESTs Weakly
25340	.2721717135	AA054554	EST
28106	.2659103748	AA485084	ESTs
38690	.2649184307	AA600121	ESTs
20203	.2626499431	N28855	ESTs Modera
10251	.2608760694	R76185	ESTs Weakly
12684	.2604192389		ESTs
31636	2509469427		Natural resista
20769	.2479765348		ESTs
1572	.2353281083		EST - K01884
10923	.2292322072		ESTs
34380			ESTs
	.2283792392		
10132	.2222816115		EST - R35733
16629	.2161752119		ESTs
25146	.1969683794		ESTs
28730	.1965943098	D20959	ESTs Modera
10200	.1874912391		ESTs
38695	.1545794663		ESTs
31365	4.150549979	N67550	ESTs
42379	.1496120668	W37999	ESTs
28050	.1428703354	AA479139	Acid phosphat
2620	.1386565707		Human recom
8927	.1340593744		Homo sapiens
13379	.1269549188		ESTs Weakly
5134	.1218251808		Human done
2626			Zinc finger pro
38005	.1160483666		ESTs
36575	.1127196584		EST
18296	.1121837207		ESTs Weakly
29531			
29531	.1111459313		EST - RC_H8
	. 1095880506	AFFX-HUMTE	ESTs
143	2222		
143 10970	.0967613396		
143 10970 25836	.0952825397	AA152305	Interferon (gah
143 10970 25836 19735	.0952825397 .0937927853	AA152305 H53038	Interferon (gar
143 10970 25836 19735 40711	.0952825397 .0937927853 .090970943	AA152305 H53038 N53564	Interferon (gar EST ESTs
143 10970 25836 19735	.0952825397 .0937927853	AA152305 H53038 N53564	Interferon (gar

FIGURE 12 (cont.)

other other other other other TM other other TM other other other other other other SS, other other other SS, SS, тм

other ? other

other other other other other other other other other other other

other other other other

TM ? other SS. ? other TM other

5503	.0861035825 X0		Stromelysin
20310	.0641711656N3		ESTs Highly
456	.0599824566 D3		Prostaglandin
7814	.0559685576 AA		ESTs
40230	.0447282719H9		ESTs
33651	4.039204804 W		ESTs
16777	.0231657929 AA		EST
19110	.0094905222 HO	8778	ESTs
34442	.0077010365 AA	258093	HKR-T1
5099	4.004992433 U7	9247	Human clone
8209	.9990473163 AA	384220	ESTs
24408	.9976586074W9	90146	ESTs
26596	.9974919787 AA		ESTs
16485	.9811264008 AA		Spleen focus
32969	.9804901745W4	12451	ESTs
27006	.9799768093 AA		ESTs Weakly
29809	.9526765967N2		EST
9596	.9440163451H9		ESTs
29024	.9377933938F0		Homo sapiens
21694	.9356365584R3		Homo sapien
	3.929998104 AA		ESTs
13207	.9143752629 AA		ESTs Highly
37865			ESTS Highly
36201	.9129828172AA		
8961	.8981160269 AF		
17444	.8927133917 AA		ESTs
25869	.8919834527 AA		ESTs Highly
24862	3.89042252 Z4		ESTs Highly
26685	3.889363206 AA		ESTs
42300	.8850230366T9		ESTs
6495	.8830844863 X9		Zinc finger pro
38604	.8828045942 AA	598803	ESTs
36358	.8826713718 AA	425756	ESTs
30560	3.873276445 N4	9284	MYB PROTO
14413	.8724466158 AA	600150	ESTs
23823	.8574824967 T9	1805	Homo sapien
38158	3.853096838 AA	487021	EST
2572	.8519747554 M2		Vascular ende
40100	.8464168967H7		Laminin recep
40258	.8462992993 HS		ESTs
20944	.8461621525N7		ESTs
20411	.8459400968 N4		Homo sapien:
10345	.8457714481AA		ESTs
31261	.8451974374N6		EST
	.8378410994 AA		ESTs
8513			
13877	.8363409835 AA		ESTs
40748	.8253562321N5		EST
14509	.8152852193AA		ESTs
10281	.8065567331R8		ESTs
25284	.8044158642 AA		ESTs Weakly
6730	.7900025129Y0		H.sapiens mR
16033	.7884592402 AF	FX-HUMIS	AFFX-HUMIS
39242	.7827164808 AA	621523	ESTs
27354	.7794760435 AA	425221	ESTs
	3.777263605 U4	9188	Human place
4552	[3,7 7 7 200003 10 7		
4552 18385	.7756199108 A	227219	Homo sapien

12752	.7671137403AA421250	(ESTs
	.7601033106W60180	ESTs
42463		ESTS
10614	.7581669016 AA037357	
867	.7459337969 D87716	Human mRNA
7608	.7336047135 AA180967	ESTs
31795	3.732738742 N80703	ESTs
35377	.7273784603 AA399453	EST - RC_AA
22828	.7243928524R98192	ESTs
25240	.7243198336 AA039713	ESTs
11008	.7197381368 AA134289	ESTs Weakly
4341	.7162349944U38545	Human ARF-a
28833	.7147818393D59787	EST - RC_D5
3750	.7121007154U09279	Collagen type
17483	.6943413512AA122147	ESTs
16854	.691520847 AA055552	ESTs Weakly
3709	.689165677 U07550	Heat shock 10
1608	.6652978422L00205	KERATIN TYP
24577	.6617721053 Z38727	Homo sapiens
31032	.6570916386 N62508	ESTs
4951	.6536195433U69546	Human RNA b
37660	.6523275307AA460225	ESTs
20418	.6495357091N49209	ESTs
27995	.6485167436 AA470155	Homo sapiens
7971	.6434397185 AA287423	ESTs
27606	3.64303453 AA443793	ESTs
24677	.6427250633Z39338	ESTs Highly
11070	.6406198277AA148521	ESTs Weakly
9328	.6356048599 D89618	Homo saplens
36826	3.634689802 AA435996	ESTs
17678	.6300045795AA134275	Human HIV1
36209	.6274694477 AA421266	ESTs Weakly
34120	.6258090412AA211615	EST
38152	.6246442011AA486737	H.sapiens mR
38463	.6184693268 AA504491	ESTs Weakly
20064	.6183699978H98653	ESTS
31256	.5992620732N66152	EST
9713	.5985228843 L44338	Homo sapiens
28622	.5768056147 D11837	ESTs
38057	.5736105703 AA481549	EST - RC AA
	.5688723791D45568	IEST - RC_AA
28763		EST - RC_AA
16996	.5680705709 AA069038	ESTs Modera
28628	.5604144617 D11888	
25804	.5442954572 AA148885	ESTs
2492	.5423964239 M22898	Tumor protein
14904	.5411970737T83389	ESTs Highly
25265	.5347588502 AA043765	H.saplens RY
13606	.5327912417 AA456437	ESTs Weakly
42307	.5318436465 T96595	EST - RC_T96
1544	3.526202414 J05068	TRANSCOBA

	FOR			
Cook	PRIMEKEY	o ol i umor ov	Accessions	Comparation
CZAS	111929	3.7	R40057	prominin (mouse)-like 1
BCX2	128790	4.0	AA291725	secreted frizzled-related protein 4
CBC2	101809	5.7	1180849	Homo saplens connexin 28 (GJB2) mRNA, complete cds
CBC1	100365	4.8	078611	mesoderm specific transcript (mouse) homolog
CBC3	102618	2.5	U65932	extracellular matrix protein 1
	134804	6.3	L37036	small inducible cytokine sublamity B (Cya-X-Cys), momber 5 (epithelial-derived neutrophil-activating peptide 78)
	104209	2.1	AB000221	arnali inducible cytokine subfamily A (Cys-Cys); member 18; pulmonary and activation-regulated
CJA8	115697	8.1	AA411502	ESTs: Weakly similar to airway trypsin-like prolease (H. saprens)
	124315	5.4	H94892	v-ral simian leukemia viral oncogene homolog A (ras related)
	109415	4.3	AA227219	Homo sapians CAGF9 mRNA: partial ods SRY (sex-determining region Y)-box 9 (campometic dysplasia; autosomal sex-reversal)
	103613	5.1	Z46629	RAB6 interacting; kinesin-tike (rabkinesin6)
	109166	6.2 5.7	AA179845 AA463725	ESTs: Weakly similar to KIAA0226 (H.sapiens)
CJA9	116176 115522	5.7 8.1	AA331393	ESTs
CGA7	125852	5.7	H09290	ESTs; Weakly similar to unknown (H.sapiens)
BCN5	112244	3.1		· ESTs
COAt	132592	5.6	AA129390	
BCN7	117280	5.4	N22107	ESTs: Moderately similar to IIII ALU SUBFAMILY SC WARNING ENTRY IIII [H.saplens]
ac.	102663	4.8	U70322	karyopherin (Importin) beta 2
COA2	104660	6.0	AA007160	ESTs .
	113702	2.4	T97307	ESTs; Moderately similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII [H.sapiens]
	100154	6.0	D14657	KIAA0101 gene product
	102260	3.7	U28386	Human nuclear localization sequence receptor hSRP taipna mRNA, complete cds
	101809	5.7	M86849	Homo sapiens connexin 26 (GJB2) mRNA, complete cds
	133272	3.2	AA465016	ESTs; Highty similar to serine protease homolog
	100365	4.8	D78611	mesoderm specific transcript (mouse) homolog
	126519	4.7	AA305536	EST176522 Colon carcinoma (Caco-2) cell line II Homo sapiens cDNA 5' end, mRNA sequence.
	132543		AA417152	ESTs; Highly similar to protein regulating cytokinesis 1 [H.sapiens]
	103023	5.5	X53793	mutifunctional polypeptide similar to SAICAR synthetase and AIR carboxylase
	132109	3.1	AA599801	ESTE
	104037	3.5	AA372630	differentially expressed in hematopoietic tineages
	104978	3.5	AA088458	ESTs; Weakly similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII [H.saplens]
	108695	3.0	AA121315	ESTs SCA
	107248	3.8 3.4	059894	ESTs ESTs
	132902	4.0	AA490969 W95477	ESTs .
	128790	4.0	AA291725	secreted frizzied-related protein 4
	101923	3.8	S75256	HNL=neurophil lipocalin [human, ovarian cancer cell line OC6, mRNA Partial, 534 nd
	119943	3.4	W86835	copine III
	130648	3.9	AA075427	EST:
	132358	3.5	X60488	H4 histone family; member G
	106286	3.2	AA434441	Irizzled (Drosophila) nomolog 7
	117557	2.3	N33920	diubiquilin
	129691	3.3	X06700	collagen; type III; alpha 1 (Ehlers-Danios syndrome type IV; autosomal dominant)
	114767	4.5	AA148885	minichromosome maintenance deficient (S. coravisiae) 4
	100335	3.8	D63391	platetel-activating factor scelythydrotase; isoform (b; gamma subunit (29kD)
	134969	3.5	AA238324	ESTs; Weakly similar to I'll ALU CLASS A WARNING ENTRY HI! [H.sapiens]
	110009	3.4	H10933	ESTs .
	124059	4.0	F13573	ESTs
	104755	2.2	AA024482	ESTs; Weakly similar to coldermal type I keratin [i4.saplens]
	107151	3.4	AAB21169	ESTS
	132669	2.9	AA185376	ESTS; Weakly senilar to 60S RIBOSOMAL PROTEIN L22 [H.sapiens]
	104394	5.5	H46617	yp19h1.r1 Soares breast 3NbM8st Homo sapiens cDNA clone IMAGE:187921 5', mRNA sequence
	117667	2.5	N39214	ser-The protein kinase related to the myotonic dystrophy protein kinase ESTs; Moderately similar to (define not available 4753768) [H.sapiens]
	104954 132994	3.3 3.7	AA074514 AA505133	ESTS WOOGERS & BURNET TO GREWING USE SANDERS AS DESTROY TO THE SERVICE OF THE SER
	102681	3.7	U72761	karyopherin (importin) bela 3
	103988	2.2	AA314779	ESTE: Weakly similar to LITHOSTATHINE I BETA PRECURSOR (H.sapiens)
	132183	2.5	L19183	Human MAC30 mRNA; 3' end
	.52.00			

FIGURE 13A

118695

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N71781
                    HG2167-HT22Protein Kinase Ht31, Camp-Dependen
100552
                    AA251829 ESTs: Moderately similar to (define not available 4680697) [H.sapiens]
120471
                    U47732
                    AA417067
125103
                    T95333
                                 ESTs; Weakly similar to Strabismus [D.melanogaster]
135243
                    AA215333
                                ESTs
                                ESTs: Moderately similar to alternatively spliced product using exon 13A [H.sapiens]
121457
                    AA411448
131216
                    D31058
                                 ESTs
112971
                    T17185
111179
                    N67239
                                 ESTs; Moderately similar to IIII ALU SUBFAMILY SC WARNING ENTRY IIII [H.sabiens]
                                 ESTs; Weakly similar to unknown (S.cerevisiae)
                                 ESTs: Moderately similar to IIII ALU SUBFAMILY SO WARNING ENTRY IIII (H.sapiens)
105156
111223
                    N68921
                                 ESTs; Weakly similar to neogenin (H.sapiens)
132180
                    AA405569
                                 fibroblast activation protein; aipha
                    AA447621 ESTs
106400
                    AA093834
                                ESTs; Highly similar to (defline not available 4679014) [H.sapiens]
129260
                    AA279943
115291
128628
                    AA599729
                                 Homa saplens homeobox protein A10 (HOXA10) gene; complete cds
                    R45698
                                ESTs: Weakly similar to Similarity to S. Pombe BEM1/BUD5 suppressor (C.olegans)
105082
                    AA143763
           3.2
                                 H.capiens mRNA for SMT3B protein
103453
                    X99585
                    AA443793
                                ESTs
115947
                                ESTs; Highly similar to (define not available 4589929) [H.sapiens]
105012
                    AA116036
                                ESTs: Moderately similar to (define not available 4106061) (H.sapiens)
105507
                    AA256678
                                 ESTs: Weakly similar to katanin p80 subunit [H.sapiens]
130800
                                ESTs: Moderately similar to IIII ALU SUBFAMILY SQ WARNING ENTRY IIII [H.sapiens]
129945
                    AA232104
                                ESTs: Highly similar to (define not available 4929579) (H.saplens)
100864
                    HG4297-HT45Transcriptional Coactivator Pc4
128131
                    AI283162
                                claudin 3
                    AA491465 EST6
131564
           2.8
100279
                    D42084
                                Human mRNA for KIAA0094 gane; partial cds
134405
                                collagen; type XI; alpha 1
108828
                    AA131584
                                ESTs; Weakly similar to coded for by C. elegans cDNA cm16f8 [C.elegans]
131289
                    AA485697
                                ESTS
109141
           4.2
                    AA 176428 ESTs
119307
           2.5
                    T32108
                                ESTS
                                ESTs; Moderately similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII [H.sapiens]
           2.1
134319
                    AA 129547
133458
                   M18728
                                non-specific cross reacting antigen
                                 ESTS; Weakly similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII (H.sapiens)
116732
```

FIGURE 13B

	- 600			
Code	PRIMEKEY	l Tumor o	Accession /	Complete Title 2019
CZAS	111929	3.7	R40057	prominin (mouse)-like 1
8CX2	128790	4.0	AA291725	secreted frizzled-related protein 4
CBC2	101809	5.7	M86849	Homo sapiens connexin 26 (GJB2) mRNA, complete cds
CBC1	100365	4.8	D78611	mesoderm specific transcript (mouse) homotog
CBC3	102618	2.5	U65932	extracellular matrix protein 1
CJA8	115697	8.1	AA411502	ESTs; Weakly similar to airway trypsin-like protease [H.saptens]
CJA9	116176	5.7	AA463725	ESTs; Weakly similar to KIAA0226 [H.sapiens]
CGA7	115522	8.1	AA331393	ESTs
BCN5	112244	3.1	R51309	ESTs (now Sulfatase by in-house sequencing) ·
CQA1	132592	5.6	AA129390	ESTs
BCN7	117280	5.4	N22107	ESTs; Moderately similar to !!!! ALU SUBFAMILY SC WARNING ENTRY !!!! [H.sapiens]
CQA2	104660	6.0	AA007160	ESTs

TSTC/STOTOGO TO A TOWN AS A SAME ATTOWN AS A SAME AT A TOWN AS A SAME AT TOWN AS A S GGGGTCAGGACTCCAGTACCTCCAAGAGAAGACTCTCTGTGGGCTTCTACTGCCATCAGCCAACGACGCTCGCAAGGGCG GTCTTGTGGCCCTGCAAGTCAGAGTACCTTGGGTCTGAAGGGGTCACTCAAGGGCTCTGCTATCTCTGCAGCTAAAACGG GTGACCGTGTCTGGGGGCACCCAAAAAGGCGAGGCTGTGCTTGGGACACACAATTAAAGACCATCACGGGGAATTCTGC TGCTGTTATTACCCCATTCAAGTTGACAACTGAGGCAACGCAGACTCCAGTCTCCAATAAGAAACCAGTGTTTGATCTTA AAGCAAGTTTGTCTCGTCCCCTCAACTATGAACCACAAAGGAAAGCTAAAACCATGGGGGCAATCTAAAGAAAATAAT ACGGAAGAAACGCGAGCAAGAACGAAAGGAAAGAAAGGAAAGGTTTTGGGAATGCGAAGGGGCCTCATTTTGGCTGAAG ATCCCCACTTAGTCACGAGATCTTTTTCTGCTAACTGTTCATAGTCTGTGTAGTGTCCATGGGTTCTTCATGTGCTATG ATCTCTGAAAAGACGTTATCACCTTAAAGCTCAAATTCTTT<u>GGGATGGTTTTACTTAAGTCCATTAACAATTCAGGTTT</u> CTAACGAGACCCATCCTAAAATTCTGTTTCTAGATTTTTAATGTCAAGTTCCCCAAGTTCCCCCTGCTGCTGATATATTA ACAGAACTGCAGTCTTCTGCTAGCCAATAGCATTTACCTGATGGCAGCTAGTTATGCAAGCTTCAGGAGAATTTGAACAA
TAACAAGAATAGGGTAAGCTGGGATAGAAAGGCCACCTCTTCACTCTCTATAGAATATAGTAACCTTTATGAAACGGGGC CATATAGTTTGGTTATGACATCAATATTTTACCTAGGTGAAATTGTTTAGGCTTATGTACCTTCGTTCAAATATCCTCAT AACTCCACCAAATGTACTTCTTATTCATTTTATGGAAAAGACTAGGCTTTGCTTAGTATCATGTCCATGTTTCCTTCACC AAAAAAAAAAAAA

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Amino-CKVLG MRRGL ILAED-COOH

FIGURE 18

Acetyl-KQPHL QTKEE QRKKC-Amide

numan_CAA2 nouse_CAA2	ARFKKMESIDEYTMRKKKHLKEHSSLNELKLDKKGIVTPVPPRGRLSVPCTPARQQCP
numan_CAA2	QGRSCGPASOSTLGLKGSLKRSAISAAKTGVRFSAATKDNEHKRSLTKTPARKSAHVTVS QG
human_CAA2	GGTQKGEAVLGTHKLKTITGNSAAVITPFKLTTEATQTPVSNKKPVFDLKASLSRPLNYE
mouse_CAA2	GSASKGQAVFRTPKSKATERTSIAVITPFKLMTEATQTPSSSKKPVFDLKASLSRPLNYK
human_CAA2	PHKGKLKPWGQSKENNYLNQHVNRINFYKKTYKQPHLQTKEEQRKKREQERKEKKAKVLG
mouse_CAA2	PHKGKLKPWGOAKENNSLNERVSRVTFHRKTYKQPHLQTREERWKRQEQERKEKKEKLLE
human_CAA2	MRRGLILAED-
mouse_CAA2	ARRNLGVTKAQ

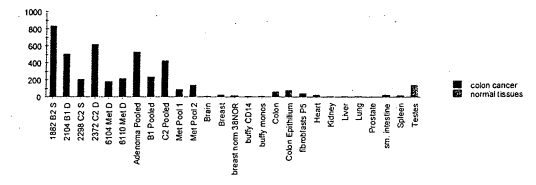


FIGURE 21

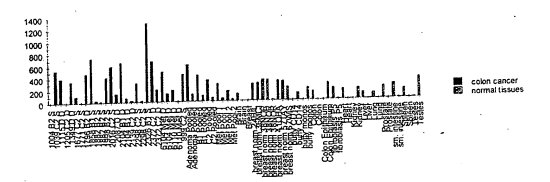
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<u>ATG</u>GAGCCGCGGCCCACGGCGCCCTCCTCCGGCGCCCCGGGACTGGCCGGGGGAGACGCCGTCAGCCGCTGCGCT GGCCGCAGCCAGGGTGGAACTGCCCGGCACGGCTGTGCCCTCGGTGCCGGAGGATGCTGCGCCGCGAGCCGGGACGGGACGGCG GCGGGGTCCGCGATGAGGGCCCCGCGGCGGCCGGGGACGGGCTGGGCAGACCCTTGGGGCCCACCCCGAGCCAGAGCCGT TGGTGCTGGGGGGGGGCCAAGCAGACCCCCGGGGCGGACGGGGAAGCCAGCGAGGGAGCCAGCTAAAGGCAGCGAGG AAGCCAAGGGCCGCTTCCGCGTGAACTTCGTGGACCCAGCTGCCTCCTCGTCGGCTGAAGACAGCCTGTCAGATGCTGCC GGGGTCGGAGTCGACGGGCCCAACGTGAGCTTCCAGAACGGGGGGACACGGTGCTGAGCGAGGGCAGCAGCCTGCACTC CGGCGGCGGCGGCGGCAGTGGGCACCACCAGCACTACTATTATGATACCCACACCCAACACCTACTACCTGCGCACCTTCG GCCACAACACCATGGACGCTGTGCCCAGGATCGATCACTACCGGCACACAGCCGCGCAGCTGGGCGAGAAGCTGCTCCGG CCTAGCCTGGCGGAGCTCCACGACGAGCTGGAAAAGGAACCTTTTGAGGATGGCTTTGCAAATGGGGAAGAAAGTACTCC AACCAGAGATGCTGTGGTCACGTATACTGCAGAAAGTAAAGGAGTCGTGAAGTTTGGCTGGATCAAGGGTGTATTAGTAC GTTGTATGTTAAACATTTGGGGTGTGATGCTTTTCATTAGATTGTCATGGATTGTGGGTCAAGCTGGAATAGGTCTATCA GTCCTTGTAATAATGATGGCCACTGTTGTGACAACTATCACAGGATTGTCTACTTCAGCAATAGCAACTAATGGATTTGT AAGAGGAGGAGGAGCATATTATTTAATATCTAGAAGTCTAGGGCCAGAATTTGGTGGTGCAATTGGTCTAATCTTCGCCT TTGCCAACGCTGTTGCAGTTGCTATGTATGTGGTTGGATTTGCAGAAACCGTGGTGGAGTTGCTTAAGGAACATTCCATA CTTATGATAGATGAAATCAATGATATCCGAATTATTGGAGCCATTACAGTCGTGATTCTTTTAGGTATCTCAGTAGCTGG AATGGAGTGGGAAGCAAAAGCTCAGATTGTTCTTTTGGTGATCCTACTTCTTGCTATTGGTGATTTCGTCATAGGAACAT TTATCCCACTGGAGAGCAAGAGCCAAAAGGGTTTTTTGGTTATAAATCTGAAATATTTAATGAGAACTTTGGGCCCGAT CTCAGGTGATCTTGCAGATCCTCAGTCAGCCATACCCAAAGGAACACTCCTAGCCATTTTAATTACTACATTGGTTTACG TAGGAATTGCAGTATCTGTAGGTTCTTGTGTTGTTCGAGATGCCACTGGAAACGTTAATGACACTATCGTAACAGAGCTA ACAAACTGTACTTCTGCAGCCTGCAAATTAAACTTTGATTTTTCATCTTGTGAAAGCAGTCCTTGTTCCTATGGCCTAAT CAGCATTAGCATCCCTAGTGAGTGCTCCCAAAATATTTCAGGCTCTATGTAAGGACAACATCTACCCAGCTTTCCAGATG TTTGCTAAAGGTTATGGGAAAAATAATGAACCTCTTCGTGGCTACATCTTAACATTCTTAATTGCACTTGGATTCATCTT AATTGCTGAACTGAATGTTATTGCACCAATTATCTCAAACTTCTTCCTTGCATCATATGCATTGATCAATTTTTCAGTAT TCCATGCATCACTTGCAAAATCTCCAGGATGGCGTCCTGCATTCAAATACTACAACATGTGGATATCACTTCTTGGAGCA ATTCTTTGTTGCATAGTAATGTTCGTCATTAACTGGTGGGCTGCATTGCTAACATATGTGATAGTCCTTGGGCTGTATAT TTATGTTACCTACAAAAAACCAGATGTGAATTGGGGATCCTCTACACAAGCCCTGACTTACCTGAATGCACTGCAGCATT CAATTCGTCTTTCTGGAGTGGAAGACCACGTGAAAAACTTTAGGCCACAGTGTCTTGTTATGACAGGTGCTCCAAACTCA CGTCCAGCTTTACTTCATCTTGTTCATGATTTCACAAAAAATGTTGGTTTGATGATCTGTGGCCATGTACATATGGGTCC TCGAAGACAAGCCATGAAAGAGATGTCCATCGATCAAGCCAAATATCAGCGATGGCTTATTAAGAACAAAATGAAGGCAT TTTATGCTCCAGTACATGCAGATGACTTGAGAGAAGGTGCACAGTATTTGATGCAGGCTGCTGGTCTTGGTCGTATGAAG CCAAACACACTTGTCCTTGGATTTAAGAAAGATTGGTTGCAAGCAGATATGAGGGATGTGGATATGTATAAACTTATT TCATGATGCTTTTGACATACAATATGGAGTAGTGGTTATTCGCCTAAAAGAAGGTCTGGATATATCTCATCTTCAAGGAC AAGAAGAATTATTGTCATCACAAGAGAAATCTCCTGGCACCAAGGATGTGGTAGTAAGTGTGGAATATAGTAAAAAGTCC GATTTAGATACTTCCAAACCACTCAGTGAAAAACCAATTACACACAAAGTTGAGGAAGAGGATGGCAAGACTGCAACTCA ACCACTGTTGAAAAAAGAATCCAAAGGCCCTATTGTGCCTTTAAATGTAGCTGACCAAAAGCTTCTTGAAGCTAGTACAC AGTTTCAGAAAAACAAGGAAAGAATACTATTGATGTCTGGTGGCTTTTTGATGATGGAGGTTTGACCTTATTGATACCT TACCTTCTGACGACCAAGAAAAAATGGAAAGACTGTAAGATCAGAGTATTCATTGGTGGAAAGATAAACAGAATAGACCA TGACCGGAGAGCGATGGCTACTTTGCTTAGCAAGTTCCGGATAGACTTTTCTGATATCATGGTTCTAGGAGATATCAATA CCAAACCAAAGAAAGAAATATTATAGCTTTTGAGGAAATCATTGAGCCATACAGACTTCATGAAGATGATAAAGAGCAA GATATTGCAGATAAAATGAAAGAAGATGAACCATGGCGAATAACAGATAATGAGCTTGAACTTTATAAGACCAAGACATA CGTGGGAATCATCAGAGTGTCCTTACCTTCTATTCATAA

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RGNHOSVLTFYS

Peptide names
CAA9p1
CAA9p2
CAA9p3
CAA9p4
CAA9p4MAPS
CAA9p5
CAA9p5MAPS

Solubility	
1mg/1ml H2O	H-CDPAASSSAEDSLSD-NH2
1mg/1ml H2O	AC-KKSDLDTSKPLSEKC-NH2
1mg/1ml H2O	AC-PLLKKESKGPIVPLC-NH2
min.amt.DMSO/H2Q	AC-EHSILMIDEIC-NH2
Img/ml buffer pH7.5	AC-EHSILMIDEIC-on 8-Branch Maps
1mg/1ml H2O	Ac-DFREEETC-NH2
	AC-DFREEETC-on 8-Branch Maps
1mg/1ml H2O	



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TAGCAGTTTGTTAACCTGACATCTCTGCCAGTCTAGTTTCTGGCCAGGTTTCCTGTCAGTAT
TCCCCCTCCTCTTTGCATTAATCAAGGTATTTGGTAGAGGTGGAATCTAAGTGTTTGTATGTC CAATTTACTTGCATATGTAAACCATTGCTGTGCCATTCAATGTTTGATGCATAATTGGAC CTTGAATCGATAAGTGTAAATACAGCTTTTGATCTGTAATGCTTTTATACAAAAGTTTATT CCCWTTAGTGAGKSWTAATTTTAGCTTGGCACTGGCCGTCGTTTTACAACGTCGTGACTGGGAAAACCCTGGCGTTACCCAACTTAATCGCCTTGCAGCACATCCCCCTTTCGCCAGCTGAATGGCGTAATAGCGAAACGCCCGCACCGCACCGCCCCTTCCCAACAGTTGCGCAGCTGAATGGCGAATGGC CTTTYCCCKCAAGCTNTAAATCGGGGC

if more than 2 bases disagree with consensus seque more than 10 positions are unkno if more than 10 positions are gap characters AI674283 j src=genbank seqtype=est dir=3' description="tz15696.x1 A1922877 | src=genbank seqtype=est dir=3' description="wn64f08.x1 医阿斯氏性抗反应性 A)979181 | src=genbank seqtype=est dir=3' description="wr71f12.x1 AA331393 | src=genbank seqtype=est dir=3' description="EST35243 ## F # # _ . AA551863 j src=genbank seqtype=est dir=3' description="ris38f08.s ****** ******* AA9087391 src=gennank seqtype=est dir=3' description="og75h07.s 医多类性性免疫性 医多种抗性毒素 AA146858 | src=genbank segtype=est dir=3' description="zo41h11.s 安徽省政立政政党员 经存货过度放送过度的第一个一个一个一个一个一 AA768709 | src=genbank seqtype=est dr=3" description="oc86e09.s 公司等有政策的 医骶右侧韧带 ------AA648921 [src=genbank seqtype=est dir=3' description="ns41b07.s AA736861 | src=genbank seqtype=est dir=3" description="ox23e09.s SECRETARY. AW070824 | src=genbank seqtype=est dir=3' description="xa30g07. ********* AI8314831 src=genbank segtype=est dir=3" description="wi49e96.x1 ------Al382409 | src=genoank seqtype=est dir=3' description="ta72a08.x1 医通常性性原体 医多种性性原因的复数形式 AIC94155 | src=genbank seqlype=est dir=3" description="qa34e09.s A1193598 | src=genbank seqtype=est dir=3' description="qe71b04.x AA775633 | src=genbank seqtype=est dir=3' description="zf2606.s AA939106 | src=genbank seqtype=est dir=3' description="oq22c12.s AA581615] src=genbank seqtype=est dir=3' description="nc84h09.s 医医生物性医胃皮肤 英名医哲学医系第一一 Al992158 | src=genbank seqtype=est dir=3' description="wi75c08.x" AI382916 | src=genbank seqtype=est dir=3' description="ic20d09.x1
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FIGURE 28A

AI874283 | src=gendank sedtype=est dir=3' description="12":5006.x1 NCI_CGAP_UI2 Home sabletts cDNA clone IMAGE:2288527 3', mRNA s' srci=gbcu7/29773 AB22677 | 3:c*genbank seqtype=est dir=3' description="wn64i08.51 NCL_CGAP_Lu19 Homo salores cDNA done IMAGE:2450247.3', InRNA "strd=gbest36/51593 A1979181 | src=genbank seqtype=est dir=3" description="wi7 1112.x1 NCI_CGAP_UI1 Homo sapiens cDNA clone IMAGE:2493167-3"; mRNA s' src!xq AA331393 | src=genbank sedtype×est dr=3" description="EST35243 Embryo, 8 week (Homo sapiens cONA 3" end, mRNA sequence," src1=gbest13/573 AA651863 | src=genbank seqtype=est dir=3" description="og/5h07.51 NCI_CGAP_GCB1 Homo sapiens cDNA done IMAGE:1185927 3", mRNA " srd=gbest17/53265 AA908739 | src=genbank seqtype=est dir=3" description="og/5h07.51 NCI_CGAP_Ox8 Homo sapiens cDNA done IMAGE:1454173 3", mRNA s" srd=gbest21/813 AA146856 | src=genbank seqtype=test dir=3" description="zo4 th 11.s1 Stratagene endotheist cell 937223 Homo sagrens cDNA done likt" srd=gbest10/20752

AA768709 | src=genbank seqtype=test dir=3" description="zo4 th 11.s1 Stratagene endotheist cell 937223 Homo sagrens cDNA done likt" srd=gbest10/20752

AA768709 | src=genbank seqtype=test dir=3" description="zo466693.51 NCI_CGAP_GCB1 Homo sagrens cDNA done IMAGE:1356616.3", mRNA " srd=gbest19/36742 AAB48921 | src=genoank seqtype=est dx=3' description='ns41b07.st NCI_CGAP_GCB1 Homo sabiens cDNA clone IMAGE:1186165 3', mRNA * srct=gbest17/50927 AA738861 { src=genbank seqtype=est dr=3" descriptions*ca23e09 st NCL CGAP_GCB1 Home salties cDNA idone IMAGE:1305832 3". mRNA " srd=gbest1870299" AW070824 { src=genbank seqtype=est dr=3" descriptions*ca23e05 1. NCL CGAP_Br18 Home salties cDNA idone IMAGE:2568346 3". mRNA " srd=gbest3861005 A/831463 (src=genbank seqtype=est dat=3' description="wy49e06.x1 NCL_CGAP_Lu19 Homo saprens cDNA done IMAGE:2406176 3', mRNA * stct=gbcst35/22416 A332409 | src-genbank seqtype=est dir=3" description="ta72a08.st Soares, total_fetus_No2HF8_9w Homo sapiens cDNA done IMAGE:20" srd=gbest28/68899
A094155 | src-genbank seqtype=est dir=3" description="ta72a08.st Soares, total_fetus_No2HF8_9w Homo sapiens cDNA done IMAGE:1688860 3", mR" srd-gbest28/16508
A094155 | src-genbank seqtype=est dir=3" description="ta73a08.st Soares, NhHMPu_S1 Homo sapiens cDNA done IMAGE:1688860 3", mR" srd-gbest28/16508 A1193598 | sic=genbank seqlype=est dir=3' description="de? 1004.x1 Source_fotal_tung_NbHL19W Homo sapiens cDNA clone IMAGE:17443" srd=gbest24/35325 AT19390 | Sickgenbarik seqtyperest oit=3 descripedhar der 1004.x1 Sobres_lotal_ning_nont.tam mono sapiens cunka cone inhabe; (1443) straffoetar/s5256

AA775633 | srchgenbarik seqtyperest dir#3' description="z/2606.s1 Sobres_tetal_near_Nbi H19W Homo sapiens cDNA clone iMAGE;3760" strafgbest1943614

AA939106 | srchgenbarik seqtyperest dir#3' description="co22c12.s1 NCI_CGAP_GC4 Homo sapiens cDNA clone IMAGE;1567094 3'; mRNA s* srchgbest21/28363 AA581615 | src=genbank seqtype=est dir=3" description="ric84h09.s1 NCI_CGAP_GC1 Homo saciens cDNA clone IMAGE:797537 3", mRNA se" srd=gbest1643395 A1992158 | src=genbank sagtype=est dir=3" description="wi75c08.x1 Soares thymus_NHFTh Homo sapiens cONA clone IMAGE:2513294 3;" src=gebest37/59215 AJ382916 | sicrgenbank seqtyperest dire3' descriptione"(c2009 xt Spares, NAHMP)_S1 Homo saptens cDNA done IMAGE:2064401 3', mR' strd=gbest26/69408

AAS86521 | src=genbank seqtyperest dire3' descriptione"(c20409.st NCt_CGAP_GC1 Homo saptens cDNA done IMAGE:797489 3', mRNA se" srd=gbest16/48294 AJT43202 | src=genbank seqtype=est dir=3" description="Ag\$9e02.x1 Soarcs_NSF_F6_9W_OT_PA_P_S1 Homo sapiens cDNA done IMAGE:237" srct=gbcu8/18434 src=genbank seqtype=est dir=3' description="zb\$3e04.s1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:30732" srcf=gbest6/55806 AA284818 | src=genbank seqtype=est dir=3' description="zt24e09.s1 Soares ovary tumor NoHOT Homo sapiens cONA clone IMAGE:714088' src1=gbest12/8311 AD46620 | src=genbank seqtype=est dir=3" description="qp4605.x1 NCL_CGAP_Co8 Homo septens cDNA done IMAGE:1926081 3", mRNA s" srd=gthest26/33599 R.C.AA465473 | src=genbank seqtype=est dir=5' describuon="aa22g08.rl NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:814046 5', mRNA s' srcf=gbest15/9675 T24696 [stc=genbank seqtype=est dir=? describion="eST473 Human coloretat cancer Homo saptens cDNA clone 17812, mRNA seque" std=gets13/6394 R.C.AA406456] stc=genbank seqtype=est dir=? describion="EST473 Human coloretat cancer Homo saptens cDNA clone 17812, mRNA seque" std=gets13/6394 R.C.AA406456] stc=genbank seqtype=est dir=? describion="2">totalent seqtype=est d R.C.AA096093 | scregenbank segrype=est dir=5' description="18199.seq.F Human total heart. Lambda ZAP Express Homo septems cDNA 5', "scrt=gbest9/36451 R.C.AA379531 | src=genbank seqtype=est dir=5' description="EST92376 Skin tumor't Homo saptems cDNA 5' end, mRNA sequence." srd=gbest13/48697 R.C.N53714 | src=genbank seqtype=est dr=5' description="yz06b05./1 Soares_multiple_sclerosis_2NbMMSP Homo sapiens cDNA done IMA" srd=gbest6/14572 H60046| src=penbank seqtype=est dir=3' description="yr19d02.s1 Soares fetal liver spleen 1NFLS Home sapiens cDNA clone IMAGE" src=gbest4/55137 RCAA77822 | scc=genbank seqtype=est dir=5" description="EST90803 Synovial sercoma Homo sapiens cDNA 5" end, mRNA sequence." srd=gbest13/46988 NS1950 | src=genbank seqtype=est dir=3' description="yx06g06.s1 Soares multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMA* srct=gbest6/12908 A372701 | src=genbank seqtype=est dir=5' description="EST64625 Colon adenocarcinoma IV Homo sapiens cDNA 5' end, mRNA sequence" src=gbest1:3/41867 Al473650 | src=genbank seqtype=est dix=3' description="tm02e10.x1 NCI_GGAP_Co14 Homo sapiens cDNA done IMAGE:2155434 3', mRNA " srd=gbest28/8505 Al225213 | scregenbank seatype=est dir=3' description="dx06g04.x1 NCI_CGAP_Lym12 Homo saptens cDNa done IMAGE:2000598.3', mRNA* srd=gbest24/66924 AA405348 | src=genbank seqtype=est dir=3' description="to/10904.s1 Soares_NhHMPu_S1 Homo sapiens cDNA done IMAGE:753198 3', mRN" srd=gbest/8/35496 | src=genbank seqtype=est dir=3' description="do/511.x1 Soares_total_tetus_Nb2HF8_9w Homo sapiens cDNA done IMAGE:19' srd=gbest/26/38475 AA749314 | src=genbank seqtype=est dn=3' description="ny12a11.s1 NCL_CGAP_GCB1 Homo septems cDNA clone IMAGE:1271516 3', mRNA " srct=gbest19/17507

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Al378205 { src=genbank seqtype=est dir=3' description="tc65a07.x1 Suares_NHMPu_S1 Homo septens cDNA clone IMAGE:2069480'3', mR* src=gbest26/64605

FIGURE 28B

METSSSDDS CDSFASDNFA NTRLQSVREG CRTRSQCRHS GFLRVAMKFP ARSTRGATNK KAESRQPSEN SVTDSNSDSE DESGMNFLEK RALNIKQNKA MLAKLMSELE SFPGSFRGRH PLFGSDSQSR RFRRRTPPGV ASRRNFERRA RELTRSRSRI LGSLDALPME EEEEEDKYML VRKRKTVDGY MNEDDLPRTR RYRSSVTLPH IIRPVEEIQK ERSWRTSAAI LEEKIITVHW ALLVINAVRR LLIPKQTAET CTAGAFEASS VAPAFETVMV KRSGMLCWIR TGIARLVEES ATAVSAGSEM DGVRLGSLCI

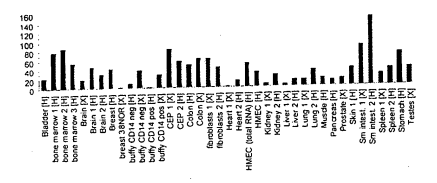


FIGURE 30A

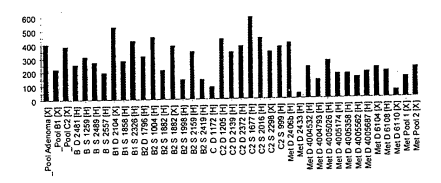


FIGURE 30B

ACTCACTATNGGGCGAATGGGCCCTCTNNATGCATGCTCGAGCGCCCGCCAGTGTGATGGATA TCTGCAGAATTCGCCCTTAAGCAGTGGTAACAACGCAGAGTACGCGGGGGGAGACCGGAGGG CAGAAGGCAGAGTCCAGGCTTAGACTGCAGTTCCTCGCTTACCTGTGCAGTCTAATTTTGAGC TGCCTCTTTGTAGTCTTAAAAGGCAGGAGCTTCGTGTTGTGGGTCTGCTAACCCGTACGTTTCC GTGGGCAAGTCGTGTGTACTCCTCGCC AAGAAATATGCCGTAGATGATGTTCCCTTCTCAATCCCTGCTGCCTCTGAAATTGCCGACCTTA GTAACATCATCAATAAACTACTAAAGGACAAAAATGAGTTCCACAAACATGTGGAGTTTGATT TCCTTATTAAGGGCCAGTTTCTGCGAATGCCCTTGGACAACACACGGAAAATGGAGAACATCT TGCATGTTCCATGATGACTGGATCAGTTCAATTAAAGGGGCAGAGGAATGGATCTTGACTGGT CATACGGATGTTGTAAAAGATGTGGCCTGGGTGAAAAAAGATAGTTTGTCCTGCTTATTATTG AGTGCTTCTATGGATCAGACTATTCTCTTATGGGAGTGGAATGTAGAGAGAAACAAAGTGAA AGCCCTACACTGCTGTAGAGGTCATGCTGGAAGTGTAGATTCTATAGCTGTTGATGGCTCAGG AACTAAATTTTGCAGTGGCTCCTGGGATAAGATGCTAAAGATCTGGTCTACAGTCCCTACAGA CTGTGGTCAGATGCTGAAGAAATCTGCAGTGCATCTTGGGACCATACAATTAGAGTGTGGGTT GTTGAGTCTGGCAGTCTTAAGTCAACTTTGACAGGAAATAAAGTGTTTAATTGTATTTCCTATT CTCCACTTTGTAAACGTTTAGCATCTGGAAGCACAGATAGGCATATCAGACTGTGGGATCCCC GAACTAAAGATGGTTCTTTGGTGTCGCTGTCCCTAACGTCACATACTGGTTGGGTGACATCAG TAAAATGGTCTCCTACCCATGAACAGCAGCTGATTTCAGGATCTTTAGATAACATTGTTAAGC TGTGGGATACAAGAAGTTGTAAGGCTCCTCTCTATGATCTGGCTGCTCATGAAGAAAAAGTTC TGAGTGTAGACTGGACAGACACAGGGCTACTTCTGAGTGGAGGAGCAGACAATAAATTGTAT TCCTACAGATATTCACCTACCACTTCCCATGTTGGGGCA TAGAGATTATTT:CTGTAAA:TGAAATTGGTAGAGAACCATGAAATTACATAGATGCAGATGCA GAAAGCAGCCTTTTGAAGTTTATATAATGTTTTCACCCTTCATAACAGCTAACGTATCACTTT TTCTTATTTTGTATTTATAATAAGATAGGTTGTGTTTATAAAATACAAACTGTGGCATACA TTCTCTATACAAACTTGAAATTAAACTGAGTITTACATTTCTCTTTAAARGTAAAAAAAAA AAAAAAAA

One position equals 17 bases. ed more than 1 bases disagree with consensus seque if more than 6 positions are unknown. d more than 6 positions are gap characters. 170 346 | 3510 (650 650 969 AW006895(13) | src=genbank seqtype=est dir=3' description="ws15g0 AA807926(0) | src=genbank seqtype=est dir=3" description="riu90e08 AA278650(15) | src=genbank seqtype=est dir=3' description="zs79a08 Al076659(17) | src=genbank seqtype=est dir=3' description="0208g04 AW058555(17) | src=genbank seqtype=est dir=3' description="wic23d0 AJ125947(18) | src=genbank segrype=est dir=3" description="qb96e12 · 表示,我们也可以是这个一个,我们们也是有一个一个,我们们就是这个一个。 AA173577(19) | src=genbank seqtype=est dir=3" description="zp04e07 AJ768162(18) | src=genbank seqtype=est dir=3' description="wh70h08 AA884295(18) | src=genbank seqtype=est dir=3' description="am15s0
AA912878(19) | src=genbank seqtype=est dir=3' description="ci27b02 AW104628(20) | src=genbank seqtype=est dir=3' description="xd84d0 AA932938(21) | src=genbank seqtype=est dir=3' description="oo06g02 · 医甲基基氏性原染性皮肤炎 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 | Al382380(21) | src=genbank seqtype=est dir=3 description="ta70h07. AJ278377(21) | src=gentrank seqtype=est dir=3' description="qm63e06 AA903844(23) | src=genbank seqtype=est dir=3' description="ok64e11 Al673354(24) | src=genbank seqtype=est dir=3' description="tw35s08. At246255(24) | src=genbank seqtype=est dir=3" description="qu39c01 Al696102(23) | src=genbank seqtype=est dire3' description="t03d10.x AA911053(23) | srumgenbank seqtypenest dir=3" description (全年表月三次為民事主法實際與公司等無法法國宣言是是正法國政治院第一一一一 AI951347(24) | src=genbank seqtype=esi dir=3" description="wx67g10 AI952510(24) | arc=genbank seqtype=est dir=3' description="wx74g05 A1417099(24) | src=genbank seqtype=est dir=3' description="1g78g12. AAS02337(26) | src=genbank seqtype=est dir=3' description="ne26a03 AA907840(32) | src=ganbank seqtype=est dir=3' description="orn16c0 AA528399(102) | src=genbank seqtype=est dir=?" description="ne83g1

FIGURE 32A

AW006895(13) | srcrogenbank seqtyperest dire3' description="ws15g01.x1 NCI_CGAP_Kid11 Homo sapiens cONA clone IMAGE:2497296 3' simil" srcrogbest37776306 AA807928(0) | src=genbank seqtype=est dir=3" description="nul90808.st NCI_CGAP_Pr22 Homo sapiens cDNA done IMAGE:1217990 3", mRNA "src=gbest1978988 AA219575(11) | src=genbank seqtype=est dir=3" description="2:05c05.st Stratagene NT2 neuronal productor 937230 Homo sapiens cDNA d" src=gbest1122843 AA27865Q(15) | src=genoank sectype=est dir=3' description="zs79a08.51 NCI_CGAP_GCB1 Homo sapiens cONA clone IMAGE:703670 3', mRNA s' srd=gbest12/4143 A076659(17) | src=genbank seqtype=est dir=3' description="roz3g04.x1 Soares_letal_tiver_spicen_1NFLS_\$1 Home sapiens cDNA done IM* srd=qbest2344980
AW058555(17) | src=genbank seqtype=est dir=3' description="roz3g04.x1 Soares_letal_tiver_spicen_1NFLS_\$1 Home sapiens cDNA done IM* srd=qbest2344980
AW058555(17) | src=genbank seqtype=est dir=3' description="roz3g04.x1 NCI_CGAP_Kd11 Home sapiens cDNA done IMAGE:2544493 3' simil* srd=qbest38/51329 Al126947(18) | src=genbank seqtype=est dir=3' description="qb96a12.x1 Soares_fetal_heart_NbHH19W Homo saptens cDNA done IMAGE:1707" srd=gbest23/50375 AA173577(19) | src-genbank seqtype=est dire3' description="xh70h08.x1 NCI_CGAP_Kid11 Homo sapiens cDNA done INAGE:2385143.3' smill* srd-gbest33/2826 AABA295(18) | src=genbank seqtype=est dir=2" description="attributed in Tunion Superior CDNA done IMAGE:1468865 3" srd=gbest2055489

AAB4295(18) | src=genbank seqtype=est dir=2" description="attributed in Tunion Superior CDNA done IMAGE:1468865 3" srd=gbest2055489

AA912876(19) | src=genbank seqtype=est dir=2" description="d27b02.s1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA done IMAGE:1524651 3" srd=gbest21/9292 AW104028(20) | strongential in sequippercest direst descriptions* x684d08.xt Soarce_NFL_T_GBC_S1 Homo sapiens cDNA done IMAGE:2604303.3* strongential in sequippercest direst descriptions* x684d08.xt Soarce_NFL_T_GBC_S1 Homo sapiens cDNA done IMAGE:2604303.3* strongential in sequippercest direst descriptions* x684d08.xt Soarce_NFL_T_GBC_S1 Homo sapiens cDNA done IMAGE:1565428.3* strongest21(26801) | strongential in sequippercest direct descriptions* x684d08.xt Soarce_NFL_T_GBC_S1 Homo sapiens cDNA done IMAGE:1565428.3* strongest21(26801) | strongest2 AS82380(21)] sro-genbank seqtype=esi dir=3' description="ta70h07.x1 Soares_total_letus_Nb2HF8_9w Homo sapiens cDNA done IMAGE:20" srd=gbest26/68381 AQ78377(21)| src=penbank seqtype=est dir=3" description="qm63e06.x1 Soares_placenta_8to9weeks_2NbHP8to9W Homo sapiens cDNA clone" srd=gbest25/38348 AB17342(23) | streightbank seqtyperest dire3' descriptions*Mx2260 xt NCI_CGAP_Lym12 Homo sapients cDNA clone IMAGE:2413044 3' similar streighests5/12499

AA903644(23) | streightbank seqtyperest dire3' descriptions*Mx2260 xt NCI_CGAP_Lym12 Homo sapients cDNA clone IMAGE:2413044 3' similar streighest21/261 AUS73354(24) | 3rc=genbank seqtype=rst dir=3' description="tw35a08.x1 NCI_CGAP_Ut1 Homo sadiens cDNA done IMAGE:2261654 3' similar' srchqbesi31/7780 AIZ46Z55(24) | src=genbank seqtype=est dir=3' description="qu39c01.x1 NCI_CGAP_Lym5 Homo saprers cDNA clone IMAGE:1967136 3' similia' srd=grest256231 ARSG102(23) 1 strongenbank settype=est dir=3' description="103d10.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2239899 3', mRNA s' strongbesi31/30525 ASS1105(23) | strongenbank seqtyperest dim3" description="old5501.31 NCL_CGAP_GC4 Homo sopiens DNA done MAGE:1518844 3' striller' stronghess1/17467
ASS0131(24) | strongenbank seqtyperest dim3" description="old5501.31 NCL_CGAP_UI3 Homo sopiens DNA done MAGE:2564539 3' striller' stronghess1/17465
ASS1147(24) | strongenbank seqtyperest dim3" description="wx67g10.x1 NCL_CGAP_Brit8 Homo sopiens DNA done MAGE:2564539 3' striller' stronghess1/174654
ASS1147(24) | strongenbank seqtyperest dim3" description="wx67g10.x1 NCL_CGAP_Brit8 Homo sopiens DNA done MAGE:2564539 3' striller' stronghess1/174616
ASS1147(24) | strongenbank seqtyperest dim3" description="wx67g10.x1 NCL_CGAP_Brit8 Homo sopiens DNA done MAGE:2564539 3' striller' stronghess1/174616
ASS1147(24) | strongenbank seqtyperest dim3" description="wx67g10.x1 NCL_CGAP_Brit8 Homo sopiens DNA done MAGE:2564539 3' striller' stronghess1/174616
ASS1147(24) | strongenbank seqtyperest dim3" description="wx67g10.x1 NCL_CGAP_Brit8 Homo sopiens DNA done MAGE:2564539 3' striller' stronghess1/174616
ASS1147(24) | strongenbank seqtyperest dim3" description="wx67g10.x1 NCL_CGAP_Brit8 Homo sopiens DNA done MAGE:2564539 3' striller' stronghess1/174616
ASS1147(24) | strongenbank seqtyperest dim3" description="wx67g10.x1 NCL_CGAP_Brit8 Homo sopiens DNA done MAGE:2564539 3' striller' stronghess1/174616
ASS1147(24) | strongenbank seqtyperest dim3" description="wx67g10.x1 NCL_CGAP_Brit8 Homo sopiens DNA done MAGE:2564539 3' striller' stronghess1/174616
ASS1147(24) | strongenbank seqtyperest dim3" description="wx67g10.x1 NCL_CGAP_Brit8 Homo sopiens DNA done MAGE:2564539 3' striller' stronghess1/174616
ASS1147(24) | strongenbank seqtyperest dim3" description="wx67g10.x1 NCL_CGAP_Brit8 Homo sopiens DNA done MAGE:2564539 3' striller' stronghess2/174616
ASS1147(24) | strongenbank seqtyperest dim3" description="wx67g10.x1 NCL_CGAP_Brit8 Homo sopiens DNA done MAGE:2564539 3' stronghess2/174616
ASS1147(24) | stronghess2/174616
ASS1147(24) | stronghess2/174616
ASS1147(24) | stronghess2/174616 A952510(24) | src-genbank seqtype=est dire3' description="tg76g12.x1 Soares_NhtMPu_S1 Homo sapiens cDNA done IMAGE:2549432 3', mRNA 'srd-gbest37/20164
A41709(24) | src-genbank seqtype=est dire3' description="tg76g12.x1 Soares_NhtMPu_S1 Homo sapiens cDNA done IMAGE:2114950 3' stm' srd-gbest27/33775 AAS02337(26) | src=genbank seqtype=est dir=3' description="ne26a03.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:898444 3', mRNA se' srd=gbest 548530 A907840(32) | sre-genbank seqtype=est dire3" description="ned3gl2.st NCL_CGAP_Ewt Homo septiens cDNA done IMAGE:1541198 3" srd-gbest21/4257

ASSE199(102) | src-genbank seqtype=est dire3" description="ned3gl2.st NCL_CGAP_Ewt Homo septiens cDNA done IMAGE:910918, mRNA seque" srd-gbest15/7439

cons for 1

FIGURE 32B

MAQLQTRFYTDNKKYAVDDVPFSIPAASEIADLSNIINKLLKDKNEFHKHVEFDF LIKGQFLRMPLDKHMEMENISSEEVVEIEYVEKYTAPQPEQCMFHDDWISSIKGA EEWILTGSYDKTSRIWSLEGKSIMTIVGHTDVVKDVAWVKKDSLSCLLLSASMD QTILLWEWNVERNKVKALHCCRGHAGSVDSIAVDGSGTKFCSGSWDKMLKIWS TVPTDEEDEMEESTNRPRKKQKTEQLGLTRTPIVTLSGHMEAVSSVLWSDAEEIC SASWDHTIRVWVVESGSLKSTLTGNKVFNCISYSPLCKRLASGSTDRHIRLWDPR TKDGSLVSLSLTSHTGWVTSVKWSPTHEQQLISGSLDNIVKLWDTRSCKAPLYDL AAHEDKVLSVDWTDTGLLLSGGADNKLYSYRYSPTTSHVGA.

FIGURE 33

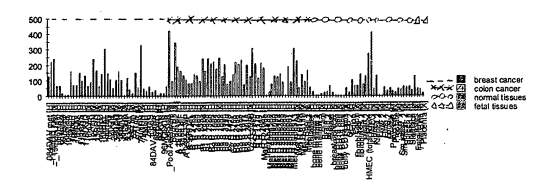


FIGURE 34

WO 00/55633 PCT/US00/07044

AGRACIA A CONTROL DE LA CACITA DE LA CACITA DE LA CACACITA DEL CACACITA DE LA CACACITA DE LA CACACITA DEL CACACITA DE LA CACACITA DEL CACACITA DE LA CACACITA DEL CACACITA DE LA CACACITA ACAGTCAGGCCCATCTGTCTGCCCTTCTTTGATGAGGAGCTCACTCCAGCCACCCCCACTCTGGATCATTG GATGGGGCTTTACGAAGCAGAATGGAGGGAAGATGTCTGACATACTGCTGCAGGCGTCAGTCCAGGTCAT TGACAGCACAGGGTGCAATGCAGACGATGCGTACCAGGGGGAAGTCACCGAGAAGATGATGTGTGCAGGC ATCCCGGAAGGGGGTGTGGACACCTGCCAGGGTGACAGTGGTGGGCCCCTGATGTACCAATCTGACCAGT GGCATGTGGTGGGCATCGTTAGCTGGGGCTATGGCTGCGGGGGCCCGAGCACCCCAGGAGTATACACCAA GGTCTCAGCCTATCTCAACTGGATCTACAATGTCTGGAAGGCTGAGCTGTAATGCTGCTGCCCCTTTGCA GTGCTGGGAGCCGCTTCCTTCCTGCCCTGCCCACCTGGGGATCCCCCAAAGTCAGACACAGAGCAAGAGT CCCCTTGGGTACACCCCTCTGCCCACAGCCTCAGCATTTGTTGGAGCAGCAAAGGGCCTCAATTCCTGTA AGCATCCCAGGGAGAGACACAGCCCACTGAACAAGGTCTCAGGGGTATTGCTAAGCCAAGAAGGAACTTT CCCACACTACTGAATGGAAGCAGGCTGTCTTGTAAAAGCCCAGATCACTGTGGGCTGGAGAGGAGAAGGA AAGGGTCTGCGCCAGCCCTGTCCGTCTTCACCCATCCCCAAGCCTACTAGAGCAAGAAACCAGTTGTAAT ATAAAATGCACTGCCCTACTGTTGGTATGACTACCGTTACCTACTGTTGTCATTGTTATTACAGCTATGG

FIGURE 35

FIGURE 36

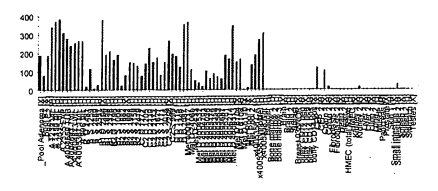


Figure 37

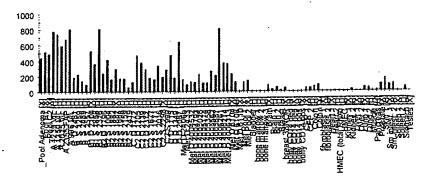


Figure 38

CCAAGTTCTACCTCATGTTTGGAGGATCTTGCTAGCT<u>ATG</u>GCCCTCGTACTCGGCTCCCTGTTGCTGCTGG GGCTGTGCGGGAACTCCTTTTCAGGAGGGCAGCCTTCATCCACAGATGCTCCTAAGGCTTGGAATTATGAA TTGCCTGCAACAAATTATGAGACCCAAGACTCCCATAAAGCTGGACCCATTGGCATTCTCTTTGAACTAGT GUNTATCTTTCTCTATGTGGTACAGCCGCGTGATTTCCCAGAAGATACTTTGAGAAAATTCTTACAGAAGG CATATGAATCCAAAATTGATTATGACAAGCCAGAAACTGTAATCTTAGGTCTAAAGATTGTCTACTATGAA GCAGGGATTATTCTATGCTGTGTCCTGGGGCTGCTGTTTATTATTCTGATGCCTCTGGTGGGGTATTTCTT TIGTATGTGTCGTTGCTGTAACAAATGTGGTGGAGAAATGCACCAGCGACAGAAGGAAAATGGGCCCTTCC TGAGGAAATGCTTTGCAATCTCCCTGTTGGTGATTTGTATAATAAGCATTGGCATCTTCTATGGTTTT GTGGCAAATCACCAGGTAAGAACCCGGATCAAAAGGAGTCGGAAACTGGCAGATAGCAATTTCAAGGACTT GUGAAUTCTCTTGAATGAAACTCCAGAGCAAATCAAATATATATTTGGCCCAGTACAACACTACCAAGGACA AGGCGTTCACAGATCTGAACAGTATCAATTCAGTGCTAGGAGGCGGAATTCTTGACCGACTGAGACCCAAC ATCATCCCTGTTCTTGATGAGATTAAGTCCATGGCAACAGCGATCAAGGAGACCAAAGAGGCGTTGGAGAA CATGANCAGCACCTTGAAGAGCTTGCACCAACAAAGTACACAGCTTAGCAGCAGTCTGACCAGCGTGAAAA AGATTGTCTCTAAGCCAGCTGAATAGCAACCCTGAACTGAGGCAGCTTCCACCCGTGGATGCAGAACTTGA CANCGTTANTANCGTTCTTAGGACAGATTTGGATGGCCTGGTCCAACAGGGCTATCAATCCCTTAATGATA TACCTGACAGAGTACAACGCCAAACCACGACTGTCGTAGCAGGTATCAAAAGGGTCTTGAATTCCATTGGT TCAGATATCGACAATGTAACTCAGCGTCTTCCTATTCAGGATATACTCTCAGCATTCTCTGTTTATGTTAA GCCTGGTCATCTGCTCTCTCCTGACCCTCATCGTGATTTTTTACTACCTGGGCTTACTGTGTGGCGTGTGC GGCTATGACAGGCATGCCACCCGACCACCGGAGGCTGTGTCTCCAACACCGGAGGCGTCTTCCTCATGGT tggagttggattaagiittcctcttttgctggatattgatgatcattgtggttcttacctttgtctttggig UNANTGTGGAAAAACTGNTCTGTGAACCTTACACGAGCAAGGNNTTATTCCGGGTTTTGGATACACCCTAC TTACTANATGANGACTGGGAATACTATCTCTCTGGGAAGCTATTTAATAAATCAAAAATGANGCTCACTTT TGANCAAGTTTACAGTGACTGCAAAAAAAATAGAGGCACTTACGGCACTCTTCACCTGCAGAACAGCTTCA ATATCAGTGAACATCTCAACATTAATGAGCATACTGGAAGCATAAGCAGTGAATTGGAAAGCTGAAGGTA nntcttnatatctttctgttgggtgcagcaggaagaaaaaccttcaggatttgctgcttgtggaataga CAGARTGAATTATGACAGCTACTTGGCTCAGACTGGTAAATCCCCCGCAGGAGTGAATCTTTTATCATTTG Catatigatetageageaaaageaaacgtttgeeeccaggaaatttgaggaaeteeetgaaaagagatgea CAAACTATTAAAACAA'ITCACCAGCAACGAGTCCTTCCTATAGAACAATCACTGAGCACTCTATACCAAAG CGTCAAGATACTTCAACGCACAGGGAATGGATTGTTGGAGAGAGTAACTAGGATTCTAGCTTCTCTGGATT TTGCTCAGAACTTCATCACAAACAATACTTCCTCTGTTATTATTGAGGAAACTAAGAAGTATGGGAGAACA ATANTAGGATATTTTGAACATTATCTGCAGTGGATCGAGTTCTCTATCAGTGAGAAAGTGGCATCGTGCAA ACCTGTGGCCACCGCTCTAGATACTGCTGTTGATGTCTTTCTGTGTAGCTACATTATCGACCCCTTGAATT "GTTTTGGTTTGGCATAGGAAAAGCTACTGTATTTTTACTTCCGGCTCTAATTTTTGCGGTAAAACTGGCT ANGTACTATCGTCGAATGGATTCGGAGGACGTGTACGATGATGTTGAAACTATACCCATGAAAATATGGA AAATGGTAATAATGGTTATCATAAAGATCATGTATATGGTATTCACAATCCTGTTATGACAAGCCCATCAC aacattgatagctgatgttgaaactgcttgagcatcaggatactcaaagtggaaaggatcacagatttttg GTAGTTTCTGGGTCTACAAGGACTTTCCAAATCCAGGAGCAACGCCAGTGGCAACGTAGTGACTCAGGCGG GCACCAAGGCAACGCACCATTGGTCTCTGGGTAGTGCTTTAAGAATGAACACAATCACGTTATAGTCCAT GGTCCATCACTATTCAAGGATGACTCCCTCCCTTCCTGTCTATTTTTGTTTTTTACACTTACACTGAGT TTCTATTTAGACACTACAACATATGGGGTGTTTGTTCCCATTGGATGCATTTCTATCAAAACTCTATCAAA GCATTTTGTGTACAGTAAACGGTGTATATACCTTTTGTTACCACAGAGTTTTTTAAACAAATGAGTATTAT AGGACTTTCTTCTAAATGAGCTAAATAAGTCACCATTGACTTCTTGGTGCTGTTGAAAATAATCCATTTTC ACTARAGEGEGEGARACCTACAGCATATECTECACGCAGAGATTTTCATCTATTATCTTTATCARAGAT TGGCCATCTTCCACTTGGAAATGGCATGCAAAAGCCATCATAGAGAAACCTGCGTAACTCCATCTGACAAA TTCANARGAGAGAGACAGATCTTGAGAGAGANATGCTGTTCGTTCAAAAGTGGAGTTGTTTTAACAGATGC CANTTACGGTGTACAGTTTAACAGAGTTTTCTGTTGCATTAGGATANACATTAATTGGAGTGCAGCTAACA TGAGTATCATCAGACTAGTATCAAGTGTTCTAAAATGAAATATGAGAAGATCCTGTCACAATTCTTAGATC TGGTGTCCAGCATGGATGAAACCTTTGAGTTTGGTCCCTAAATTTGCATGAAAGCACAAGGTAAATATTCA TTTGCTTCAGGAGTTTCATGTTGGATCTGTCATTATCAAAAGTGATCAGCAATGAAGAACTGGTCGGACAA AATTTAACGTTGATGTAATGGAATTCCAGATGTAGGCATTCCCCCAGGTCTTTTCATGTGCAGATTGCAG TTCTGATTCATTTGAATAAAAAGGAACTTGG

FIGURE 40

CAGUGGCCGCTGAATTCTAGGGCGGGTTCGCGCCCCGAAGGCTGAGAGCTGGCGCTGCTCGTGCCCTGTG "GCCAGACGGCGGACCTCCGCGGCCGGACCCCGCGGCCCCGCTTTGCTGCCGACTGGAGTTTGGGGGAAG ANACTOTUCTGCGCCCAGAAGATTTCTTCCTCGGCGAAGGCACAGCGAAAGATGAGGGTGGCAGGAAGA GANGGCGCTTTCTGTCTGCCGGGGTCGCAGCGCGAGAGGGCAGTGCCATGTTCCTCCTACTAGTGGC TGCCGGCACATGCCCTGGAACATCACGCGGATGCCCCAACCACGTGCACCACGCAGGAGAACGCCA TCCTGGCCATCGAGCAGTACGAGGAGCTGGTGGACGTGAACTGCAGCGCCGTGCTGCGCTTCTTCTTCTG TGCCATGTACGCGCCCATTTGCACCCTGGAGTTCCTGCACGACCCTATCAAGCCGTGCAAGTCGGTGTGC CAACGCGCGCGCGACGACTGCGAGCCCCTCATGAAGATGTACAACCACAGCTGGCCCGAAAGCCTGGCCT GCGNCGAGCTGCCTGTCTATGACCGTGGCGTGTGCATTTCGCCTGAAGCCATCGTCACGGACCTCCCGGA GGATGTTAAGTGGATAGACATCACACCAGACATGATGGTACAGGAAAGGCCTCTTGATGTTGACTGTAAA CGCCTAAGCCCCGATCGGTGCAAGTGTAAAAAGGTGAAGCCAACTTTGGCAACGTATCTCAGCAAAAACT ACAGCTATGTTATTCATGCCAAAATAAAAGCTGTGCAGAGGAGTGGCTGCAATGAGGTCACAACGGTGGT GGATGTNANAGAGNTCTTCNNGTCCTCATCACCCNTCCCTCGNACTCAAGTCCCGCTCNTTACAAATTCT TCTTGCCAGTGTCCACACATCCTGCCCCATCAACATGTTCTCATCATGTGTTACGAGTGGCGTTCAAGGA "GATGCT"CTTGAAAATTGCTTAGTTGAAAAATGGAGAGATCAGCTTAGTAAAAGATCCATACAGTGGGA AGAGAGGCTGCAGGAACAGCGGAGAACAGTTCAGGACAAGAAGAAAACAGCCGGGCGCACCAGTCGTAGT NATCCCCCAAACCAAAGGGNAAGCCTCCTGCTCCCAAACCAGCCAGTCCCAAGAAGAACATTAAAACTA GGAGTGCCCAGAAGAGAACAAACCCGAAAAGAGTG<u>TGA</u>GCTAACTAGTTTCCAAAGCGGAGACTTCCGAC TICCTTACAGGATGAGGCTGGGCATTGCCTGGGACAGCCTATGTAAGGCCATGTGCCCCTAACA ACTCACTGCAGTGCTCTTCATAGACACATCTTGCAGCATTTTTCTTAAGGCTATGCTTCAGTTTTTCTTT CTANGCCATCACAAGCCATAGTGGTAGGTTTGCCCTTTGGTACAGAAGGTGAGTTAAAGCTGGTGGAAAA GGCTTATTGCATTGCATTCAGAGTAACCTGTGTGCATACTCTAGAAGAGTAGGGAAAATAATGCTTGTTA CANTTEGACCTANTATGTGCATTGTANANTAAATAATGCCATATTTCAAACAAAAAAACACGTAATTTTTTTACAG TGTTTTTAAGAAGGAACAGTAGTGGAATGAATGTTAAAAAGATCTTTATGTGTTTTATGGTCTGCAGAAGGA TTTTTGTGATGAAAGGGGATTTTTTGAAAAATTAGAGAAGTAGCATATGGAAAATTATAATGTGTTTTTT ANABAGGAGAGGCAGACAATGTCTGGATTCCTGTTTTTTGGTTACCTGATTTCCATGATCATGATGCTTC TTGTCAACACCCTCTTAAGCAGCACCAGAAACAGTGAGTT1GTCTGTACCATTAGGAGTTAGGTACTAAT TAGTTGGCTAATGCTCAAGTATTTTATACCCACAAGAGGTATGTCACTCATCTTACTTCCCAGGACAT CCACCCTGAGAATAATTTGACAAGCTTAAAAAYGGCCTTCAYGTGAGTGCCAAATTTTGTTTTTCTTCAT TTAAATATTTTCTTTGCCTAAATACATGTGAGAGGAGTTAAATATAAATGTACAGAGGAAAGTTGAGT TTTGCTAACACAGTAAGCATGTATTTTATAAGGCATTCAATAAATGCACAACGCCCAAAGGAAATAAAAT CCTATCTAATCCTACTCTCCACTACACAGAGGTAATCACTATTAGTATTTTTGGCATATTATTCTCCAGGT GTTTGCTTATGCACTTATAAAATGATTTGAACAAATAAAACTAGGAACCTGTATACATGTGTTTCATAAC CTGCCTCCTTTGCTTGGCCCTTTATTGAGATAAGTTTTCCTGTCAAGAAAGCAGAAACCATCTCATTTCT AACAGCTGTGTTATATTCCATAGTATGCATTACTCAACAAACTGTTGTGCTATTGGATACTTAGGTGGTT TCTTCACTGACAATACTGAATAAACATCTCACCGGAATTC

GATTTAATCCTATGACAAACTAAGTTGGTTCTGTCTTCACCTGTTTTGGTGAGGTTGTGTAAGAGTTGGT GTTTGCTCAGGAAGAGATTTAAGCATGCTTGCTTACCCAGACTCAGAGAAGTCTCCCTGTTCTGTCCTAG CTATGTTCCTGTGTTGTGTGCATTCGTCTTTTCCAGAGCAAACCGCCCAGAGTAGAAGATGGATTGGGGC ACGCTGCAGACGATCCTGGGGGGTGTGAACAACACTCCACCAGCATTGGAAAGATCTGGCTCACCGTCC TCTTCATTTTTCGCATTATGATCCTCGTTGTGGCTGCAAAGGAGGTGTGGGGAGATGAGCAGGCCGACTT TGTCTGCAACACCCTGCAGCCAGGCTGCAAGAACGTGTGCTACGATCACTACTTCCCCATCTCCCACATC CGGCTATCGGCCCTGCAGCTGATCTTCGTGTCCAGCCCAGCGCTCCTAGTGGCCATGCACGTGGCCTACC GGAGACATGAGAAGAAGAAGTTCATCAAGGGGGAGATAAAGAGTGAATTTAAGGACATCGAGGAGAT CANANCCCHGANGGTCCGCATCGAAGGCTCCCTGTGGTGGACCTACACAAGCAGCATCTTCTTCCGGGTC ATCTTCGAAGCCGCCTTCATGTACGTCTTCTATGTCATGTACGACGGCTTCTCCATGCAGCGCTGGTGA RGTGCAACGCCTGGCCTTGTCCCAACACTGTGGACTGCTTTGTGTCCCGGCCCACGGAGAAGACTGTCTT CACAGTGTTCATGATTGCAGTGTCTGGAATTTGCATCCTGCTGAATGTCACTGAATTGTGTTATTTGCTA ATTNGATATTGTTCTGGGAAGTCNAAAAGCCAGTTTAACGCATTGCCCAGTTGTTAGATTNAGAAATNG ACAGCATGAGAGGGATGAGGCAACCCGTGCTCAGCTGTCAAGGCTCAGTCGCCAGCATTTCCCAACACAA AGATTCTGACCTTAAATGCAACCATTTGAAACCCCTGTAGGCCTCAGGTGAAACTCCAGATGCCACAATG TTAGTTCCACTGAGACCCCAGGCTGTTAGGGGTTATTGGTGTAAGGTACTTTCATATTTTAAACAGAGA TATCGGCATTTGTTTCTTTCTCTGAGGACAAGAGAAAAAGCCAGGTTCCACAGAGGACACAGAGAAAGGT TTGGGTGTCCTCCTGGGGTTCTTTTGCCAACTT1CCCCACG1TAAAGGTGAACATTGGTTCTTTCAT1T GC"TTGGNNGTTTTNNTCTCTANCAGTGGACAAAGTTNCCAGTGCCTTAAACTCTGTTACACTTTTTGGA AGTGAAAACTTTGTAGTATGATAGGTTATTTTGATGTAAAGATGTTCTGGATACCATTATATGTTCCCCC TGTTTCAGAGGCTCAGATTGTAAYATGTAAATGGTATGTCATTCGCTACTATGATTTAATTTGAAATATG ACCTAACAACATTGTAGCCTCAATCGAGTGAGACAGACTAGAAGTTCCTAGTTGGCTTATGATAGCAAAT GGCCTCATGTCAAATATTAGATGTAATTTTGTGTAAGAAATACAGACTGGATGTACCACCAACTACTACC TGTAATGACAGGCCTGTCCAACACATCTCCCTTTTCCATGCTGTGGTAGCCAGCATCGGAAAGAACGCTG **NTTTAAAGAGGTGAGCTTGGGAATTTTATTGACACAGTACCATTTAATGGGGAGACAAAAATGGGGGCCA** GGGGAGGGAGGTTTCTGTCGTTAAAACGAGTTTGGAAAGACTGGACTCTAAATTCTGTTGATTAAAG NTGACCTTTGTCTACCTTCAAAAGTTTGTTTGGCTTACCCCCTTCAGCCTCCAATTTTTTAAGTGAAANT ATAACTAATAACATGTGAAAAGAATAGAAGCTAAGGTTTAGATAAATATTGAGCAGATCTATAGGAAGAT TGAACCTGAATATTGCCATTATGCTTGACATGGTTTCCAAAAAATGGTACTCCACATACTTCAGTGAGGG TAAGTATTTTCCTGTTGTCAAGAATAGCATTGTAAAAGCATTTTGTAATAATAAAGAATAGCTTTAATGA TATGCTTGTAACTAAAATAATTTTGTAATGTATCAAATACATTAAAACATTAAAATATAATCTCTATAA

GAGTGGCTGTAGCTGCCCCGGCGGCGGCTGCCGCCCTGCGGGCTGTGGGCTGCGGGCTGCGCCCCCGCT GCATANCGCGCCCATGGTGCGCCGAGATCGCCTCCGCAGGATGAGGGAGTGGTGGGTCCAGGTGGGGCTG CTGGCCGTGCCCTGCTTGCTGCGTACCTGCACATCCCACCCCTCAGCTCTCCCCTGCCCTTCACTCAT CCANGTCTTCAGGCAAGTTTTTCACTTACAAGGCACTGCGTATCTTCTACCAAGACTCTGTGGGTGTGGT TGGANGTCCAGAGATAGTTGTGCTTTTACACGGTTTTCCAACATCCAGCTACGACTGGTACAAGATTTGG AAGCGTCTGACCTTGAGGTTTCATCGGGTGATTGCCCTTGATTTCTTAGGCTTTGGCTTCAGTGACAAAC CGAGACCACATCACTATTCCATATTGAGCAGGCCAGCATCGTGGAAGCGCTTTTGCGGCATCTGGGGCT CCNGARCOGCAGATCARCCTTCTTTCTCATGACTATGGAGATATTGTTGCTCAGGAGCTTCTCTACAGG AGACTUACUGTUCACTCUTTUTCUAAAGCTACTUAAGATGGAGGTGTGUTGTCACCCATCUTCACACG ACTGATGAACTTCTTTGTATTCTCTCGAGGTCTCACCCCAGTCTTTGGGCCGTATACTCGGCCCTCTGAG AGTGAGCTGTGGGACNTGTGGGCAGGGATCCGCAACNATGACGGGNACTTAGTCATTGNCAGTCTCTTAC TCATTTTATCTATGGGCCATTGGATCCTGTAAATCCCTATCCAGAGTTTTTGGAGCTGTACAGGAAAACG CTGCCGCGGTCCACAGTGTCGATTCTGGATGACCACATTAGCCACTATCCACAGCTAGAGGATCCCATGG GCTTCTTGAATGCATATATGGGCTTCATCAACTCCTTCTTATGGCTGGAAAGAGTAGCTTCCCTGTATTACC TCCCCTACTCCCTTATGTGTTGTGTATTCCACTTAGGAAGAATGCCCAAAAGAGGTCCTGGCCATCAAA CATATTCTCTCACAAAGTCCACTTTACTCAAATTGGTGAACAGTGTATAGGAAGAAGCCAGCAGGAGCT CTGACTAAGGTTGACATAATAGTCCACCTCCCATTACTTTGATATCTGATCAAATGTATAGACTTGGCTT TGTTTTTTGTGCTATTAGGAAATTCTGATGAGCATTACTATTCACTGATGCAGAAAGACGTTCTTTTGCA TANAAGACTTTTTTTTAACACTTTGGACTTCTCTGAAATATTTAGAAGTGCTAATTTCTGGCCCACCCCC ANCAGGAATTCTATAGTAAGGAGGAGAGAGGGGGGGCTCCTTCCCTCTCGAATGACGTTATGGGCA CATGCCTTTTAAAAGTTCTTTAAGCAACACAGAGCTGAGTCCTCTTTGTCATACCTTTGGATTTAGTGTT TCATCAGCTGTTTTTAGTTATAAACATTTTGTTAAAATAGATATTGGTTTAAATGATACAGTATTTTAGG ACTCTGCCAGAGTGGAGCTAATTAAACACGTTTGGTTTCTGAATAAATTGAACTAAATCCAAACTAT TTCCTAAAATCACAGGACATTAAGGACCAATAGCATCTGTGCCAGAGATGTACTGTTATTAGCTGGGAAG ACCANTTCTANCAGCAAATAACAGTCTGAGACTCCTCATACCTCAGTGGTTAGAAGCATGTCTCTTGA CCTACAGTAGAGGGGAAGGGATTGTTGTGTAGTCAAGTCACCATGCTGAATGTACACTGATTCCTTTATG ATGACTGCTTAACTCCCCACTGCCTGTCCCAGAGAGGCTTTCCAATGTAGCTCAGTAATTCCTGTTACTT TACAGACAGGAAAGTTCCAGAAACTTTAAGAACAAACTCTGAAAGACCTATGAGCAAATGGTGCTGAATA CTTTTTTTTTTAAAGCCACATTICATTGTCTTAGTCAAAGCAGGATTATTAAAGTGATTATTTAAAATTCGT TTTTTTAAATTAGCAACTTCAAGTATAACAACTTTGAAACTGGAATAAGTGTTTATTTTCTATTAAAA AATGAATTGTGACAAAAAAAAACCG

CTCTGAGTGTCCAGTGGTCAGTTGCCCCAGGATGGGGACCACAGCCAGAGCAGCCTTGGTCTTGACCTAT TTGGCTGTTGCTTCTGCTGCCTCTGAGGGAGGCTTCACGGCTACAGGACAGAGGCAGCTGAGGCCAGAGC ACTITCANGAAGTTGGCTNCGCAGCTCCCCCTCCCCACCCTATCCCGAAGCCTCCCCATGGATCACCC ACCCCTCTCCAACAGGAAAAGCTGCTACCTGCCCAACTCCCTGCTGAAAAGGAAGTGGGTCCCCTCTCC GCC/AGCTCCATTTGGGGACCAGAGCCATCCAGAACCTGAGTCCTGGAATGCAGCCCAGCACTGCCAACAG CACCGGTCCCAAGGGGGCTGGGGCCACCGCTGGATGGCTTCCCCCTGGGCGGCCTTCTCCAGACAATC TGNACCNNNYCTGCCTTCCTAACCGTCAGCNTGTGGTATATGGTCCCTGGNACCTACCACAGTCCAGCTN CTCCCACCTCACTCGCCAGGGTGAGACCCTCAATTTCCTGGAGATTGGATATTCCCGCTGCTGCCACTGC CGCACCCACACAAACGCCTAGAGTGTGCCAAACTTGTGTGGGAGGAAGCAATGAGCCGATTCTGTGAGG CCAGGAGGAAGCTCCCCAGCCACACTACCAGCTCCGGGCCTGCCCCAGCCATCAGCCTGATATTTCCTCG **GUTCTTGAGCTCCCTTTCCCTCCTGGGGTGCCCACATTGGACAATATCAAGAACATCTGCCACCTGAGGC** CCTTCCGCTCTGTGCCACCCAACCTGCCAGCTACTGACCCCCTACAAAGGGAGCTGCTGGCACTGATCCA GCTGGAGGGGAGTTCCAGCGCTGCTGCCGCCAGGGGAACAATCACACCTGTACATGGAAGGCCTGGGAG GATACCCTTGACAAATACTGTGACCGGGAGTATGCTGTGAAGACCCACCACCACTTGTGTTGCCGCCACC CTCCCAGCCCTACTCGGGATGAGTGCTTTGCCCGTCGGGCTCCTTACCCCAACTATGACCGGGACATCTT GACCATTGACATCAGTCGAGTCACCCCCAACCTCATGGGCCACCTCTGTGGAAACCAAAGAGTTCTCACC AAGCATAAACATATTCCTGGGCTGATCCACAACATGACTGCCCGCTGCTGTGACCTGCCATTTCCAGAAC AGGCCTGCTGTGCAGAGGAGAAATTAACCTTCATCAATGAYCTGTGTGCCCCGACGTAACATCTG GCGAGACCCTGCCCTCTGCTGTTACCTGAGTCCTGGGGATGAACAGGTCAACTGCTTCAACATCAATTAT CTGAGGAACGTGGCTCTAGTGTCTGGAGACACTGAGAACGCCAAGGGCCAGGGGAGCAGGGCTCAACTG GAGGAACAAATATCAGCTCCACCTCTGAGCCCAAGGAAGAATAAGTCACCCCAGAGCCCTAGAGGGTCAG

ACTUACTATAGGGCTCGAGCGGCCGGCCGGGCAGGTGGCCACCACCATCATCTAAAGAAGATAAACTTGG CANATGACATGCAGGTTCTTCAAGGCAGAATAATTGCAGAAAATCTTCAAAGGACCCTATCTGCAGATGTT CTGAATACCTCTGAGAATAGAGATTGATTATTCAACCAGGATACCTAATTCAAGAACTCCAGAAATCAGGA GACGGAGACATTTTGTCAGTTTTGCAACATTGGACCAAATACAATGCAAGTATTCTTGCTGTGCTCTGGTTT TGGCTGTCCTGGGCACAGAATTGCTGGGAAGCCTCTGTTCGACTGTCAGATCCCCGAGGTTCAGAGGACGG ATACAGCAGGAACGAAAAACATCCGACCCAACATTATTCTTGTGCTTACCGATGATCAAGATGTGGAGCT CGGGTCCCTGCAAGTCATGAACAAAGGAGAAAGATTATGGAACATGGGGGGGCCCACCTTCATCAAYGCCT TTGTGACTACACCCATGTGCTGCCCGTCACGGTCCTCCATGCTCACCGGGAAGTATGTGCACAATCACAAT ATATOTTANCANCACTGGCTACAGAACAGCCTTTTTTGGAAAATACCTCAATGAATATAATGGCAGCTACA TCCUCCCTGGGTGGCGAGAATGGCTTGGATTAATCAAGAATTCTCGCTTCTATAATTACACTGTTTGTCGC ANTGCCATCANAGAAAAGCATGGATT"GATTATGCAAAGGACTACTTCACAGACTTAATCACTAACGAGAG CATTANTTACTTCAAAATGTCTAAGAGAATGTATCCCCATAGGCCCGTTATGATGGTGATCAGCCACGCTG CGCCCACGGCCCGAGGACTCAGCCCCACAGTTTTCTAAACTGTACCCCAATGCTTCCCAACACATAACT CCTACTTATAACTATGCACCAAATATGGATAAACACTGGATTATGCAGTACACAGGACCAATGCTGCCCAT CCNCNTGGAATTTACANACATTCTACAGCGCAANAGGCTCCAGACTTTGATGTCAGTGGATGATTCTGTGG GGTTACCATATTGGGCAGTTTGGACTGGTCAAGGGGAAATCCATGCCATATGACTTTGATATTCGTGTGCC TTTTTTTTTTCGTGGTCCAAGTGTAGAACCAGGATCAATAGTCCCACAGATCGTTCTCAACATTGACTTGG CCCCACGATCCTGGATATTGCTGGGCTCGACACACCTCCTGATGTGGACGGCAAGTCTGTCCTCAAACTT AGTGGAAAGGCAAATTTCTACGTAAGAAGGAAGAATCCAGCAAGAATATCCAACAGTCAAATCACTTGC CCAAATATGAACGGGTCAAAGAACTATGCCAGCCAGGCCAGGTACCAGACAGCCTGTGAACAACCGGGGCAG AAGTGGCAATCCATTGAGGATACATCTGGCAAGCTTCGAATTCACAAGTGTAAAGGACCCAGTGACCTGCT CACAGTCCGGCAGAGCACGCGGAACCTCTACGCTCGCGGCTTCCATGACAAAGACAAAGAGTGCAGTTGTA CCAAAGTACAAGCCCAGATTTGTCCATACTCGGCAGACACGTTCCTTGTCCGTCGAATTTGAAGGTGAAAT ATATGACATAAATCTGGAAGAAGAAGAAGAAGTGCAAGTGTTGCAACCAAGAAACATTGCTAAGCGTCATG ATGAAGGCCACAAGGGGCCAAGAGATCTCCAGGCTTCCAGTGGTGGCAACAGGGGCAGGATGCTGGCAGAT AGCAGCAACGCCGTGGGCCCACCTACCACTGTCCGAGTGACACAAGTGTTTTATTCTTCCCAATGACTC TATCCATTGTGAGAGAGACTGTACCAATCGGCCAGAGCGTGGAAGGACCATAAGGCATACATTGACAAAG AGATTGAAGCTCTGCAAGATAAAATTAAGAATTTAAGAGAAGTGAGAGGACATCTGAAGAGAAGAAGAAGCCT GAGGAATGTAGCTGCAGTAAACAAAGCTATTACAATAAAGAGAAAGGTGTAAAAAAAGCAAGAGAAATTAAA GAGCCATCTTCACCCATTCAAGGAGGCTGCTCAGGAAGTAGGAAACTGCAACTTTTCAAGGAGAACA CTCACTTGCTTCACGCATGACAACAACCACTGGCAGACAGCCCCGTTCTGGAACCTGGGATCTTTCTGTGC TTGCACGAGTTCTAACAATAACACCTACTGGTGTTTGCGTACAGTTAATGAGACGCATAATTTTCTTTTCT GTGAGTTTGCTACTGGCTTTTTGGAGTATTTTGATATGAATACAGATCCTTATCAGCTCACAAATACAGTG CACACGGTAGAACGAGGCATTTTGAATCAGCTACACGTACAACTAATGGAGCTCAGAAGCTGTCAAGGATA TANGCAGTGCAACCCANGACCTAAGAATCTTGATGTTGGAAATAANGATGGAGGAAGCTATGACCTACACA GAGGACAGTTATGGGATGGATGGGAAGGTTÄÄTCAGCCCCGTCTCACTGCAGACATCAACTGGCAAGGCCT AGAGGAGCTACACAGTGTGAATGAAACATCTATGAGTACAGACAAAACTACAGACTTAGTCTGGTGGACT GGNCTARTTACTTGAAGGATTTAGATAGAGTATTTGCACTGCTGAAGAGTCACTATGAGCAAAATAAAACA AATAAGACTCAAACTGCTCAAAGTGACGGGTTCTTGGTTGTCTCTGCTGAGCACGCTGTGTCAATGGAGAT CGCCTCTGCTGACTCAGATGAAGACCCAAGGCATAAGGTTGGGAAAACACCTCATTTGACCTTGCCAGCTG ACCTTCAAACCCTGCATTTGAACCGACCAACATTAAGTCCAGAGAGTAAACTTGAATGGAATAACGACATT CCAGAAGTTAATCATTTGAATTCTGAACACTGGAGAAAAACCGAAAAATGGACGGGGCATGAAGACTAA TCATCTGGAAACCGATTCAGTGGCGATGGCATGACAGAGCTAGAGCTCGGGCCCAGCCCCAGGCTGCAGC CCATTCGCAGGCACCCGAAAGAACTTCCCCAGTATGGTGGTCCTGGAAAGGACATTTTTTGAAGATCAACTA TATUTTCCTGTGCATTCCGATGGAATTTCAGTTCATCAGATGTTCACCATGGCCACCGCAGAACACCGAAG TAATTCCAGCATAGCGGGGAAGATGTTGACCAAGGTGGAGAAGAAATCACGAAAAGGAGAAGTCACACCACC TAGAAGGCAGCGCCTCCTCACTCTCCTCTCATTAGATGAAACTGTTACCTTACCCTAAACACAGTATT TCTTTTTNNCTTTTTTATTTGTAAACTNNTNNNGGTAATCACAGCCACCAACATTCCAAGCTNCCCTGGGT ACCTTGTGCAGTAGAAGCTAGTGAGCATGTGAGCAAGCGGTGTGCACACGGAGACTCATCGTTATAATTT ACTATCTGCCAAGAGTAGAAAGAAAGGCTGGGGATATTTGGGTTGGCTTGGTTTTGATTTTTTGCTTGTTT GTTTGTTTTGTACTAAAACAGTATTATCTTTTGAATATCGTAGGGACATAAGTATATACATGTTATCCAAT

FIGURE 45A

CANGNIGGCTAGAATGGIGCCTTTCIGAGTGICTAAAACTIGACACCCCTGGTAAAICTTTCAACACACTT CCACTGCCTGCGTAATGAAGTTTTGATTCATTTTTAACCACTGGAATTTTTCAATGCCGTCATTTTCAGTT TTATCAGTCTCACTGTTGGCTGTCATTGTGACAAGTCAAAGTCAAATAAACCCCCAAGGACGACACACAGTATGGA CCACATATTGTTTGACATTAAGCTTTTGCCAGAAAATGTTGCATGTGTTTTACCTCGACTTGCTAAAATCG ATTITCTTGACAGGAGAAAAAGATCTAAAGATCTTTTATTTTCATCTTTTTTGGTTTTCTTGGCATGACTA AGAAGCTTAAATGTTGATAAAATATGACTAGTTTTGAATTACACCAAGAACTTCTCAATAAAAGAAAATC ATGAATGCTCCACAATTCCAACATACCACAAGAGAAGTTAATTTCTTAACATTGTGTTCTATGATTATTTG IAAGACCTTCACCAAGTTCTGATATCTTTTAAAGACATAGTTCAAAATTGCTTTTGAAAATCTGTATTCTT GAAAATATCCTTGTTGTGTATTAGGTTTTTAAATACCAGCTAAAGGATTACCTCACTGAGTCATCAGTACC CTCCTATTCAGCTCCCCAAGATGATGTGTTTTTGCTTACCCTAAGAGAGGTTTTCTTCTTATTTTTAGATA GTTATAGCTGAATCTTTTGGTAACTTTAAATCTTTATCATAGACTCTGTACATATGTTCAAATTAGCTGC PTGCCTGATGTGTATCATCGGTGGGATGACAGAACAAACATATTTATGATCATGAATAATGTGCTTTGT AAAAAGATTTCAAGTTATTAGGAAGCATACTCTGTTTTTTAATCATGTATAATATTCCATGATACTTTTAT AGAACAATTCTGGCTTCAGGAAAGTCTAGAAGCAATATTTCTTCAAATAAAAGGTGTTTAAAACTTTAAAAA **ΑΑΑΑΑΛΛΑΑΑΑΛΛΑ**ΑΛΛΛ

FIGURE 45B

CANGANTICGGCACGAGGGCGTGTTCCNGAAGTGCTGGATACTGTCAGTAGTTAATTCTCAGCTGGCAGGT TCCCTGAGTGCAGCTGGCTCGATAGTCGTAAATGAAGAGTGTGTCCGAAAAGACTTTGAATCCAGTATGAA TGTAGTACAGGAAATTAAATTTAAGTCTAGGATCAGAGGGACTGAAGACTGGGCTCCTCCTAGATTTCAAA TCATATTTAATATTCATCCACCACTCAAGAGGGACCTTGTGGTGGCAGCCCAGAATTTTTTCTGTGCCGGC TGTGGAACTCCAGTAGAGCCTAAGTTTGTGAAGCGGCTCCGGTACTGCGAATACCTAGGGAAGTATTTCTG TGACTGCTGCCACTCATATGCAGAGTCGTGCATCCCTGCCCGAATCCTGATGATGTGGGGACTTCAAGAAGT ACTACGTCAGCAATTTCTCCAAACAGCTGCTCGACAGCATTATGGCACCAGCCCATTTTCAATTTGCTGAGC ATCGGCCANAGCCTGTATGCGAAAGCCAAGGAAGCTGGACAGAGTGAAGGAAATTCAGGAGCAGCTCTTCCA TATCANAGAAGCTGTTGAAGACCTGTAGGTTTGCTAMCAGTGCATTAAAGGAGTTCGAGCAGGTGCCGGGA CACTTGACTGATGACTCCACCTGTCCTCCCTTAGGACCTGGTCAGGATCAAGAAAGGGCTGCTGGCACCTT NTTTGTGAATTTTCCCAGAATACGACTGTCATCTTCCCATTTCAGACAGCAACATGTAGAAGATGTTCAGC GTRCAGGGCTTGCTTTCACAAACAGTGCTTCCAGTCCTCCGAGTGCCCCGGTGTGCGAGGATCACAGCGA GGAGAAAACTTCTGGAAAGTGTGGCCTCTGCAGCAACATGATGCCCCTGAGTACTGTGAAAAAAGACTGTTC AACATGCCTTATGATAACACCGATTTGTGTCTATTATTGGTGACATTGTTTTAGATATTGGGTATTGTATA TTANGGAAAAGATGGTCTATATTCTCTTTATTGCATATACTTAATGTTTCAAAAGAATGCAGATTCTGTG TTTAAGCACAGGGCTGATAGTTGTGTGTTTTGCAAATGTTCTGTTTTGGCTGCTATTGGTTTTTAAA TATATGTGAACTGTAACTGACAAGATGAATTACTCAGTTTCTCTTTCTCTAAAGCTTGTTTGATGAAACTG GTTGGTCCTTTCNAGTGMAMAAAWATATGACCCCAAAAAAAAAAAAAAAAAAAAAAAAAGCAATGCGAAGGTG CTNATCCTGACGCTCGTGGTGGCCGCCTGCGGCTTCGTCCTCTGGAGCAGCAATGGGCGACAAAAGAA C:GAGGCCCTCGCCCCACCG1"TGCTGGACGCCGAACCCGCG

FIGURE 46

AATTTTTGGCCCCTCCGAAGGCCCAAAGAAATTTCCGGCCACCGAAGGAATTTTTGGTACCACCAGGGGG Granatggaartgcttttcaaacttagtttcctttccatttcttagtctggcctttgacacaaat CTGGTAGAAAGAAGCCTGATAAATTGAGGGCASTTGTACCCTCCTGTGCCCCAGAAGGTTCTTGGAGAG CTGCTCACCTTGCTGGGAGCTTTATTCTGATCTCATTTTGATGTTCCAGAGGGAGCATCATAAGAGCCCA CAGCTCCGATTTCCAAAGAGTGATATTGACATTTATGGAGATTGGTGTTGTAACATATTTTGATAAATACT CGTTATTTTCTACATAGGGAAAGAGAATTCGAGGGATAGACAGTCTCCAAGAAAAGTGAAGTGGTGGGAGA CCAATAGACTATTAGAGTTGACATTTGACATTTIAATGGGCGCCATGGCTCATTTTGTAGATTGAGAAGG TGCCTCTCCCCTGCTCCAAGTCTCATGATGACAGCGTGCTGACAGCTGGGAGTCTGTGGCCTTCCTCACGC AGAGGCCTTAAAGCTGGACACAGAAGCACGCCTAGGCTGGGCAGGGATGGGACCCATGCCCCCTCCTTAGA GGACGGGCTTCCTGGTTAGGAAAGGACACGTGGGGGGGGCCTTGCATAATAGTTCACTGGTCACCGTGCTTT TATGAGTAGTGTTTTTGTGCACTTGCCAGGGGTTTTCTCTCTGTGTGCGAGGGGAGTGATTTAAGCAATGG AANTACACACACAAAAAAAAAAAATGATAGTTTCACATCTCTTAGTTCCCTTGCCCAAACAAGAATATTCT "attgfgtctactttatctgtgcaccagccacaatacccacattggaaagacccatttgtgatgggtaa ACATCCCTTCCTGTCTCCCACAACCCCTGTGACTGCCCTGCATGTGTTCATGACCTCCGAAGGCCCTAATT CATGAAGCAGCAAACCCAGCAGATCTCCACCCCCTGCCTCAGGACCTCTGCTGAAGAGGGGGGATGAAGTG GGTCTCCAGGGAGGCAGTGGGGGCCTTGTTGGCAGCTGGCTCGGGAGCCGGCTTACAGGAGGGCAGCTCTG CAGTTGGGAGGGGCACCGTCCGGAGGAGACCAGGCCTCTACACACCCCCCACTCTACTTATCATCCCTGCT TTCCCCAGGCTGTCTCAGAAAGGTCGCTAAATGTATACTGTTGTCAGAATTGCTGAGATCTCCCCCCACT TCCCTTGTGCTGACTCTAGCACAGTGGCCAGGATCCAATACGAGTCCAGGGGTGACCGCAGGATGGTGGGG GCAGCGGCTTCTCCACCTACCCCAGCCACCAAGGCCCTGACGCACTTCCCTCCACCCTTCAGCACATCC CTGTGCACAGCTGGAAGGGTGCATGGCCCGCTCACCTTTGTTCAGATGGGTGGAAACGCTGATGATACCAG TGUCCTCUAGCCCAACCCACAGTGGAGACTGGAGACAGGGCAATGAGTCTGGTCGGGGGCACGTGGACATG CCCCATAGGGGCCCCACCCAGACTTAACAGGCAAGGTCCTGGGCATTGCGCGACGCAGGACTCAATGCTAA ACCANGCCTGCCTGTCTGTGCCAGGGCCCCTCTTCTGATTTACACATCCCATTTTTACACAGACCCTTC CTTCTTAATAAAGGCTGACAGTTCTGTTGGCAGCCAAGAACCCACACCATGAAGACAGGGAGTGAGGGGCC TTTGTGCCCAACTCCAGCACACCTGCGTTCTGGGGTGTGTGAGAGGCATGTTCGTGTCTGTGCGCTGGTGG TCTCGTGAGACAGTTCCGAGGACGGGGAAATTGCAGGGTGGTGGGGGCGTGAGGCTTATATGTGGAACTGA TGCAGAGTTCGCCTGCAGACGGATCTGGATATACACTATGTATAATTGTTACGTGTAATTTAAAATATATC TGTTTGCCATCGTCATGAGAAGATTATATGTAAGGCTCTGAAGGGAGAGGGAGATGTACATTCTGCCAGGC TCCTGGGGACCTTATCCGAGTCATGAAATTGATGACTGTTGATCCAGTGGTGCAAGAAGCTACACTCCATG TGTCATCACGCTTATGACTCCTAATGTATTTTTAAGGCAAAAAATGTCAGCCGACTCCATCTTCACCCCTC GATTCCTCGAGTCCAGCCTTTCTGTGCCAGTGCTTCACTGAGCCACAACGCTCTCGCCATCGGGACCCGGC TGGGCCTGGAGTCTGGGGGCACAGTTGCCATGGAGCCCTCCTGGGTCATTCTACAAATGTGCTGAGTGCCA GTGTGTCCAGCAGGGTCAGGAAGCAGGATGGAAAGATCCATTCAGACTGTTAATTTATTAACAAGGCAAAT GATTTTGTGTTTTCTTGATGACAGACTATTAAGTTTGGGACTTATTTTCCCATTTGAGAAGTTATAATATAT ATTTAAGATGATAAGTTTCCTGCTTAAGTTGTGCCTTTCAGCTTCAATGAGTTTAAGGAGCACTAAGGGTA ATGATACCARTGAGGGTTGGTTTATTATCAAACCTGAATAGCTGTGGTTTCTCCAGTAAATATTTTCTTCT TGATGTTCTATTTTCTAATAGTTTTCTTTAGTTTCTTAAAGTTGTGATACTAGATTTAGATTCTGATGC TAACTECAAATCAGGTTGGTCTCTGCTGGGTCTCTCCTGCTTTTATTTTACTTTAAGGACAAGTGTAGTTG ACCACTTCCTACCATACTTTATGTTGTAAAATCAAACTCTTTTGTGGTACATTATCTCATGCTTCTGCAAA

FIGURE 48

genome IE-35

490 JABO16492.1 Homo sapiens hJTB gene, complete cds e-I 18 491 X98176 H.sapiens mRNA for MACH-beta- I protein IE-36 Homo sapiens huntingtin interacting protein

HYPK mRNA,

492 AF049613 partial cds 7E-22

493 AF039690.1, HomosapiensantigenNY-CO-8(NY-CO-8) mRNA, partialeds IE-37

INM-001003IHomo sapiens ribosomal protein, large, PI ribosomal

494 phosphoprotein PI mRNA, complete cds. 4E-3. . .